

Spectral Line Shapes in Astrophysics and Related Topics

Edited by Milan S. Dimitrijević and Luka Č. Popović Printed Edition of the Special Issue Published in *Atoms*



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Special Issue Editors Milan S. Dimitrijević Luka Č. Popović

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About the Special Issue Editors

Milan S. Dimitrijević (PhD) is a retired research professor at the Belgrade Astronomical Observatory. His other roles include Director of Belgrade Astronomical Observatory (1994–2002), Editor-in-Chief of the *Serbian Astronomical Journal* (1987–2002), President of the Astronomical Society "Rudjer Boskovic" (1982–2004), Editor-in-Chief of the journal *Vasiona* (Universe) (1985-2004 and from 2018), President of the Society of Astronomers of Serbia (2008–2014), Federal Minister for Science, Technology and Development (1993–1994). Member elect of the Association of Writers of Serbia (2008). Member of the Governing Board of Euro-Asian Astronomical Society, Member of the Council of European Astronomical Society (2008–2012). Since 2009, he has been an Affiliated Fellow of the Paris Observatory. He is the author of around 300 papers in international scientific journals and of more than ten books, and has worked on about 10 TV series on astronomy. His main scientific interests are the following: Stark broadening, line profiles, line broadening, astronomical spectroscopy, atomic and molecular collision processes in stellar atmospheres, astroinformatics, history and philosophy of astronomy, astronomy in culture.

Luka Č. Popović (Prof.) is a research professor at the Belgrade Astronomical Observatory, and a professor at the University of Belgrade (Faculty of Mathematics) and University of Banja Luka (Faculty of Science). His main scientific interests include active galactic nuclei, gravitational lensing, astronomical spectroscopy, and the history of science. He has published more than 400 scientific papers, 120 of which are in leading astronomical journals, and has acted as editor of a number of conference proceedings and special issues of journals.

Preface to "Spectral Line Shapes in Astrophysics and Related Topics"

The analysis of spectral lines from various astrophysical sources is a powerful tool to collect data on the various properties of objects, from the solar system to the most distant quasars. Conditions of the astrophysical plasma, and the temperatures and densities of different particles are within much wider ranges than laboratory plasma; therefore, the lines from X-ray (Fe K) to the radio domains (radio recombination lines) have been observed. For example, on the basis of observed spectra, one can perform diagnostics, analysis, and modelling of various astrophysical objects from interstellar clouds of molecular hydrogen (with an electron temperature of around 50 K and electron densities of several electrons per cubic centimeter) to neutron star atmospheres with temperatures up to ten millions of Kelvin and extremely high electron densities. Such spectra can also be used to determine the chemical composition of stellar atmospheres, determine opacities, and even obtain information on thermonuclear processes in stellar interiors. Moreover, Doppler broadening can give us information about the kinematical properties of emission gas.

To properly investigate spectral lines from cosmic sources, astronomers need the corresponding reliable atomic and molecular data and precise laboratory measurements of spectral line properties. Therefore, interactions between astronomers and physicists who investigate spectral lines and use the results of such investigations for the analysis and synthesis of spectra emitted from astrophysical/laboratory plasma are necessary. A convenient method to stimulate such interaction is to bring together astronomers and physicists in order to review and discuss the present stage of investigation, with the objective of better understanding the significance of emission/absorption line shapes for different astrophysical research topics and to improve our knowledge in this domain. With such an objective, we organized a series of conferences dedicated to considering spectral line shapes.

The 1st Yugoslav Conference on Spectral Line Shapes (I YuCSLS) has been organized by the Co-Chairmen of the Scientific Committee (SC) at the Astronomical Observatory: M.S. Dimitrijević and L. Č. Popović in Krivaja, 11–14 September, 1995. The idea was to have a conference on this topic in Serbia, in the years where there is no International Conference on Spectral Line Shapes (ICSLS), and to give young researchers an opportunity to present their results and discuss them with leading international scientists in this field. The proceedings of this conference were published in *Publications of Astronomical Observatory of Belgrade* No. 50 [1], containing eight invited lectures and contributed papers. The II YuCSLS was organized by the Faculty of Physics (Chairman J. Purić) in Bela Crkva, 29 September–2. October, 1997. Four invited lectures and 29 contributed papers are published as "Proceedings" [2], in *Publications of Astronomical Observatory of Belgrade* No. 57.

The third was organized by the Faculty of Sciences in Novi Sad (Chairman S. Djurović) and was held in Brankovac, Fruška Gora, 4–6 October, 1999. About 40 participants presented five invited lectures and 25 contributed papers, published in the *Journal of Research in Physics* (Vol. 28, No. 3) [3]. The fourth, which changed its name to the Serbian Conference on Spectral Line Shapes (SCSLS), was organized by the Belgrade Astronomical Observatory (Co-Chairmen M. S. Dimitrijević, L. Č. Popović) in Arandjelovac, 10–15 October, 2003. The proceedings, with 17 invited lectures and 16 contributed papers are published in *Publ. Astron. Obs. Belgrade* No. 76 [4]. This was the first conference with wider international participation (Russia, Greece, Italy, Tunisia, and France). After this conference, we realized that the principal group interested in this series of conferences is astronomers, so we added "Astrophysics" to the conference name. All future conferences were organized by Belgrade

Astronomical Observatory.

The V Serbian Conference on Spectral Line Shapes in Astrophysics (V SCSLSA - Co-Chairmen M. S. Dimitrijević, L. Č. Popović), was held in Vršac, 6–10 June, 2005. The proceedings are published in *Memorie della Societa Astronomica Italiana Supplementi* Vol. 7 [5]. They contain 51 papers divided into three topics: (i) spectral line phenomena in extragalactic objects; (ii) stellar and interstellar spectral lines; (iii) spectral lines in laboratory plasma.

VI SCSLSA (Co-Chairmen M. S. Dimitrijević, L. Č. Popović) was in Sremski Karlovci,11–15 June, 2007 (Proceedings were published [6] with 49 articles in *AIP Conference Proceedings*, Vol. 938).

VII SCSLSA (Co-Chairmen M. S. Dimitrijević, L. Č. Popović) was in Zrenjanin, 15–19 June, 2009. Proceedings are published in *New Astronomy Reviews*, Vol. 53 [7]. Articles (28) are divided into the following topics: (i) spectral line phenomena in extragalactic objects; (ii) stellar and interstellar spectral lines: (iii) spectral lines in laboratory plasma—Astrophysical applications.

VIII SCSLSA (Co-Chairmen L. Č. Popović, D. Jevremović) was held in Divčibare, 6–10 June, 2011. (Proceedings with 46 papers are published in *Baltic Astronomy* vol. 20 [8], and proceedings of the special session "Spectral Lines and Black Holes" [9], with five articles, in *New Astronomy Reviews*, Vol. 56).

IX SCSLSA (Co-Chairmen L. Č. Popović, M. S. Dimitrijević) was organized in Banja Koviljača, 13–17 May, 2013. Selected papers are published in the special issue of *Advances in Space Research*: "Spectral Line Shapes in Astrophysics and Related Phenomena", Vol. 54 [10]. The issue has been divided into the following sections: (i) Laboratory Astrophysics—Spectral Lines and Atomic Processes, (ii) Spectral Lines in Solar and Stellar Spectra, (iii) Extragalactic Astrophysics and Spectral Lines). A special session, "Spectral Lines and Polarization," was organized during the conference within the frame of COST action MP1104 "Polarisation as a tool to study the Solar System and beyond."

X SCSLSA (Co-Chairmen L. Č. Popović, M. S. Dimitrijević) was organized in Srebrno Jezero, Serbia, 15–19 June 2015. The conference hosted 69 participants from Algeria, Austria, Bulgaria, Croatia, France, Greece, Israel, Italy, Japan, New Zealand, Poland, Republic of Srpska (Bosnia and Herzegovina), Russia, Saudi Arabia, Serbia, Spain, Tunisia, U.K., Ukraine, and the USA. At the conference, 27 invited lectures, 19 progress reports and 32 posters were presented. Two special sessions were organized: "Line Shifts in Astrophysics" and "Spectral Lines and Compact Stars" within the frame of the COST Action MP 1304 "Exploring Fundamental Physics with Compact Stars." Selected papers were published as a special issue with 23 articles in *Journal of Astrophysics and Astronomy* (vol. 36) [11].

The following is XI SCSLSA, (Chairman Luka Č. Popović), Šabac, Serbia, 21–25 August 2017. The conference hosted 66 participants from Algeria, Brasil, Chile, China, Croatia, France, Germany, Greece, India, Israel, Italy, Republic of Srpska (Bosnia and Herzegovina), Russia, Saudi Arabia, Serbia, Tunisia, Turkey, U.K., Ukraine, and presented 15 invited lectures, 26 progress reports and 24 posters. The two special sessions presented were "Line shapes in astrophysics and fusion plasma research: Common challenges" and "Collisions and spectral line shapes (in honour of Milan S. Dimitrijević's 70th birthday)." From 65 contributions presented at the Conference, 25 selected papers were published as a special issue of the journal *Atoms* [12].

In addition to published proceedings and special issues, other materials from the mentioned eleven conferences, like photos and PowerPoint presentations, are available online in the Serbian Virtual Observatory (http://servo.aob.rs/eeditions/SCSLSA.php).

Here, papers from the Special Issue "Spectral Line Shapes in Astrophysics and Related Topics" are organized as a book and reprinted. We divided them into four groups: 1. Stark broadening data for astrophysical and laboratory plasma investigations; 2. applications of spectral lines for astrophysical and laboratory plasma research; 3. spectral line phenomena in extragalactic objects; and 4. laboratory astrophysics results for spectra investigation. In the first group are seven papers, including a review titled "Stark broadening from impact theory to simulations." It offers new quantum mechanical and semi-classical calculations of Stark broadening parameters for Ar VII lines, investigation of Stark widths of Ar II spectral lines in the atmospheres of subdwarf B stars, and new Stark broadening parameters for Se IV, Sn IV, Sb IV, Te IV and Na IV spectral lines. Additionally, the regularities and systematic trends of Zr IV Stark widths have been discussed and investigated.

The second group, "Applications of spectral lines for astrophysical and laboratory plasma research," also contains seven papers. The Stark and Zeeman effects on spectral lines in magnetized white dwarf atmospheres and tokamak edge plasmas are considered, as well as line shape modeling for the diagnostic of the electron density in a coronal discharge, and the contribution of Lienard–Wiechert potential to the electron-impact broadening of spectral lines in plasmas. Also, the effect of turbulence on shapes in astrophysical and fusion plasmas is considered and a method to determine gas temperature in argon non-thermal plasmas at atmospheric pressure is presented. Finally, the Doppler broadening of spectral lines in relativistic plasmas is investigated.

The third group "Spectral line phenomena in extragalactic objects" has four papers dealing with Active Galactic Nuclei. The search for such objects is discussed, as well as their multifrequency properties, characterization by models of emission-line profiles and spectral energy distribution, quasar black hole mass estimates from high-ionization lines and the photometric and polarimetric interpretation of the behaviour of a blasar.

The fourth group, containing seven papers, deals with laboratory astrophysics results for spectra investigations. Some collisional processes in stellar atmospheres, screening characteristics of the dense astrophysical plasmas and nonlinear spectroscopy of alkali atoms in cold medium of astrophysical relevance have been considered. Also, databases with radiative and collisional molecular data, as well as electron-scattering cross-sections of relevance for the Rosetta mission in the BEAMDB database, have been presented and discussed. Finally, the application of cut-off Coulomb potential for the calculation of bound-bound state transitions is demonstrated, and the results of the investigation of the electric discharge plasma between copper electrodes immersed into water were presented.

The reviews and research papers, relating to new research on the covered topics, presented in this book, are of interest for specialists and PhD students. We hope that this series of conferences, as well as this book, will be useful and interesting for scientists interested in the investigation of spectral line shapes and will contribute to the education of young researchers and PhD students.

> Milan S. Dimitrijević, Luka Č. Popović Special Issue Editors





Editorial Collaboration between Paris and Belgrade Observatoriesnin Scientific Researches on Spectral Line Shapes[†]

Sylvie Sahal-Bréchot

LERMA, Observatoire de Paris, PSL Research University, CNRS, Sorbonne Universités, UPMC Univ. Paris 06, 92190 Meudon, France; sylvie.sahal-brechot@obspm.fr † For the 70th anniversary of Milan S. Dimitrijević.

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In the following, I will review my scientific cooperation with Milan Dimitrijević which has lasted for almost 40 years. The cooperations of Milan with the other laboratories of the Paris Observatory are outside of the present topic.

The subject of our cooperation concerns Stark broadening and shifts of "isolated" spectral lines of neutral and ionized atoms (neighbouring levels of the line do not overlap, electrons and positive ions of the medium perturb the lines) within the impact semiclassical perturbation theory.

I developed this theory for my thesis that I defended in 1969, and I made a Fortran code for the calculations, named SCP. The results provide line widths and shifts as a function of temperature and electron density of the medium. We improved the code and exploited it in our cooperation. The work is intended to spectroscopic diagnostics and modelling in astrophysics (especially stellar physics), laboratory and technological plasmas (magnetic confinement and Tokamaks, Laser and ion-beam fusion, discharge lamps and lighting...). The achieved results have led to more than 150 publications in refereed journals, to many invited papers, oral presentations and posters, in international and national Conferences and workshops, and to more than 1500 citations in ADS for the whole work. Furthermore, we have created an online database in free access which collects our results and refers to our publications.

This cooperation was born in 1978, thanks to Prof. Petar Grujić, who wrote a letter to Henri van Regemorter, who was the director of our laboratory DAMAP (Paris Observatory, Meudon, France), and another letter to me. Milan wished to spend some time in our laboratory, to work on the theory of line broadening.

In fact, after my thesis, I only applied my SCP code to a few cases intended to spectroscopic diagnostics of astrophysical interest, and I was also moving my researches towards atomic polarization and collisional depolarization, in view of spectropolarimetric diagnostics of magnetic fields in the solar corona and prominences.

So, I was very interested by Petar Grujić and Milan's wish, and I proposed a subject of comparison between quantum close-coupling electron collisions and SCP calculations, for the case of the resonance line of neutral lithium broadened and shifted by electron collisions. Moreover, new recent experiments were just made in Belgrade on this subject. Nicole Feautrier, also from our laboratory, had a copy of a close-coupling code suitable for electron-lithium collisions. Milan came two times to Meudon-DAMAP in 1978, then several times more, and the first paper of our beginning cooperation was published in *J. Phys. B* (1981, 14, 2259–2568).Then Milan came several times to Meudon, and our cooperation developed. In fact the eighties showed a wide expansion of line broadening researches, both theoretical and experimental, especially on Stark broadening. At the same time, the needs in atomic data for stellar spectroscopic diagnostics were rapidly increasing. I gave all my computer codes to Milan, and he had the good idea to couple them to the Bates and Damgaard subroutine for obtaining the necessary atomic structure, and to enter a list of energy levels, identified by quantum numbers, taken from the existing literature. This permitted to obtain results for many lines in a same run, for a series of temperatures

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and one density only for the first version of this new code. The two founding papers (*JQSRT* **1984**, *31*, 313 and *A&A* **1984**, *136*, 289–298) marked the beginning of a numerous series of papers and of active participations to many Conferences and workshops, (invited talks, progress reports, posters...). The first one was a poster at 7th ICSLS, Aussois, France, 11–15 June 1984). In the following years, several visits of Milan followed, and led to a number of publications, especially concerning neutral simple atoms (one electron above a closed shell).

At the end of the eighties, the astrophysical interest moved towards the modelling of stellar interiors and subphotospheric envelopes, and to observations in all the domains of wavelengths. The developments of powerful computers permitted to more accurately interpret the spectra of stellar atmospheres. Plasma laboratory research also showed a great expansion. All that needed extensive sets of atomic data, including Stark broadening. In addition, the interest for databases emerged at the 1994 IAU General Assembly.

For satisfying these needs, the SCP code required improvements and updates. For instance, concerning the infrared magnesium lines observed in the solar chromosphere, transitions between high levels were better treated with the improved version. Furthermore, especially for the lines of highly charged ions, observed in stellar UV spectra, and for the modelling of stellar interiors, the SCP code needed other improvements. They were completed with an active and substantial participation of Véronique Bommier, also from my laboratory.

Then the updated SCP code could be exploited at a large scale. However, the great lists of data resulting from our calculations caused some difficulties for publishing them in refereed journals, and I had to spend a certain time in exchanging letters with one editor for defending the importance of publishing long tables. In fact, at that time, only hard-copies journals existed. Paper was expensive and long tables took a lot of space. Fortunately, Internet was growing, and ftp was born. Thus, our long tables could be published on line and downloaded. But only by those who had an Internet access! So, in addition, our tables were also published in hard copies in Bulletin de l'Observatoire Astronomique de Belgrade (BOAB).

Then, due to the war, our cooperation could only go on via Internet, e-mails and Faxes. It was impossible for Milan to travel and visit our laboratory in France until the beginning of the new century.

The beginning of the new century was marked (from my point of view) by two important evolutions:

- Growing of interdisciplinary research between nuclear physics and astrophysics, especially in view of understanding the creation of heavy (heavier than Iron) and rare earths elements by neutron capture and not by nucleosynthesis. So, accurate spectroscopic diagnostics for determining abundances were needed, since the interesting lines are weak.
- 2. Increasing development of Virtual Observatories and Databases, due to the great amount of data which needed to be connected. Interoperable e-infrastructures were urgently needed.

It is in this context that Milan could come again in 2003 in our laboratory, which became a part of a broader laboratory in Paris Observatory two years ago, named the LERMA. Numerous visits of Milan followed until now, one or sometimes two per year.

In this time, the Virtual Observatory and the creation of databases became a priority of Paris Observatory. VO-PADC (Virtual Observatory Paris Data Center) was created for organizing databases and services. The LERMA recruited engineers and technicians, and Marie-Lise Dubernet was recruited as a research scientist Astronomer by a transfer from Franche-Comté University, (France).

The LERMA encouraged us and helped us to put all our numerous tables of results in a database. It made available a technician in 2005 for scanning the oldest results, then an engineer (Nicolas Moreau) for all the technical aspects (construction and development of the database and website).

So, STARK-B (http://stark-b.obspm.fr) was born and opened on line at fall 2008 as part of the databases of LERMA-Paris Observatory, with a link to the Serbian Virtual Observatory (http://servo.aob.rs//).Today, all our SCP published data are implemented, and the implementation of all MSE (Modified-Semi-Empirical) Serbian data will be shortly finished.

In 2010, due to a huge effort of Marie-Lise Dubernet, the European project VAMDC (Virtual Atomic and Molecular Data Center - http://www.vamdc.eu/) was born, and this helped us a lot.

STARK-B is a node of VAMDC. It follows the standards of VAMDC and Virtual Observatories. Milan and myself are the responsible scientists, and Nicolas Moreau is responsible for all technical aspects. The database has been presented in a number of International and National Conferences.

The cooperation was enriched by the collaboration with Tunisia. In fact, in the nineties, Prof. Zohra ben Lakhdar, from Tunis University, sent me Nabil Ben Nessib. So, he came to DAMAP-Meudon for completing his thesis on Stark broadening and shifting. He applied the SCP theory to some oxygen lines. He also extended the semiclassical collision functions to non ideal plasmas. Just after that, Neïla Larbi-Terzi, from the same Tunis laboratory, came to DAMAP-Meudon for also completing her thesis, and she applied the SCP theory to some infrared helium lines of interest for hot early-type stars.

Then Nabil obtained a position of Professor at Carthage University, Tunis, and could recruite young research students who prepared a thesis on collisional Stark broadening. Actually, Nabil has also cooperated with Milan since 1989, and this led to several presentations in scientific meetings and conferences. Then the two cooperations Tunisia-LERMA and Tunisia-AOB merged around the middle of this century. In particular, the participations to the Serbian Conferences on Spectral Line Shapes in Astrophysics (SCSLSA) contributed to reinforce the links. In fact, Nabil's coworkers and students (now seniors!), joined the cooperation. One of them, Haykel Elabidi has just obtained a position of Professor at Bizerte. Haykel developed for his thesis a quantum code for obtaining widths for lines of ionized atoms (Superstructure and Distorted wave for the treatment od the collision) and applied it to a number of ions. Hedia ben Chaouacha developed new expressions of the semi-classical collisional functions for strongly correlated plasmas and applied them to some neutral isolated helium lines. Rafik Hamdi coupled for his thesis the SCP code to the accurate Cowan atomic structure code, very useful for complex neutral and ionized atoms and for obtaining a great number of data in a same run. He applied it to several ions. The thesis of Besma Zmerli concerned neutral copper, important for technological applications. Especially, Walid Mahmoudi developed for his thesis useful expressions intended to very complex atoms. In addition, Nabil coupled the SCP code to TOPBase (R-matrix accurate atomic structure code), useful also for obtaining a great number of data in a same run. It was applied with Neïla to C II lines of interest for white dwarfs. All these works led to a number of papers, it is impossible to cite all them there. Moreover, Nabil created essential fitting formulae as a function of temperature for STARK-B, which were urgently required by stellar physics modelling.

Several Serbian AOB scientists also efficiently participate in our cooperation: Andjelka Kovačević, Nenad Milovanović, Zoran Simić, as well as Magdalena Christova, from Bulgaria (who is now professor at Sofia). Numerous publications and participations in scientific meetings and Conferences reflect their interest and involvement in the subject.

And now, what will be the future?

In the short and medium term, we plan to:

- carry on new calculations of line widths and shifts for needs in modeling and spectroscopic diagnostics, and implementation of the resulting published data in STARK-B,
- implement our published quantum results in STARK-B,
- develop other fitting formulae (regularities, systematic trends) for STARK-B,
- improve the calculation of the shift.

In the long-term, we hope that the new generation, scientists and their students, present and to come, who will replace us, will be interested to modernize and update methods and codes.

Thank you, Milan, for this extraordinarily fruitful cooperation, which led to a so large number of publications and to the STARK-B database. I hope that it will continue as long as possible.



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Article



Stark Broadening from Impact Theory to Simulations

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Abstract: Impact approximation is widely used for calculating Stark broadening in a plasma. We review its main features and different types of models that make use of it. We discuss recent developments, in particular a quantum approach used for both the emitter and the perturbers. Numerical simulations are a useful tool for gaining insight into the mechanisms at play in impact-broadening conditions. Our simple model allows the integration of the Schrödinger equation for an emitter submitted to a fluctuating electric field. We show how we can approach the impact results, and how we can investigate conditions beyond the impact approximation. The simple concepts developed in impact and simulation approaches enable the analysis of complex problems such as the effect of plasma rogue waves on hydrogen spectra.

Keywords: stark broadening; impact approximation; numerical simulation

1. Introduction

Stark profiles are used in astrophysics and other kinds of plasmas for obtaining information on the charged environment of the emitting particles. Using light for conveying information on the plasma often requires a modeling of both the plasma and the radiator. We will review different situations requiring different modeling approaches. The impact-broadening approach considers the emitter-plasma interaction as a sequence of brief separate collisions decorrelating the radiative dipole. Impact models are very effective for many types of plasmas, and can be applied to different kinds of emitters, hydrogen being an exception for most plasma conditions. Many different models using impact approximation have been developed, and we will review the most commonly-used. One can distinguish firstly between models keeping the quantum character of the perturbers, and those using a classical trajectory for the charged particles. Full quantum approaches require specific calculation techniques, which, once established, have proved to be of general interest. Another way to look at the models is the degree of accuracy required. It is often not necessary to have an accuracy better than about 20%, since the experimental errors are often of the same order or worse. This has enabled the development of a semi-empirical impact model, useful especially in cases where one does not have a sufficient set of atomic data for adequate application of more sophisticated methods with which one can readily obtain a large number of line shapes [1], making it an effective diagnostic tool.

A typical starting point for a line shape formalism in a plasma is a full quantum formalism for the emitter and the perturbers. It can be written as a linear response for the emitter dipole operator, and

provides the response of an emitter at a time t, knowing its state at an initial time [2]. This response in time allows the physical measurement of the spectrum to which it is linked by a Fourier transform. Quantum formalism introduces specific computational difficulties, but also brings powerful tools such as the angular averages. We will briefly discuss such approaches, and how they compare to semi-classical calculations. Classical path impact approximations have been widely developed, and exist in several levels of accuracy, depending on whether one is interested in a rapid analysis of a large number of spectra, or one asks for an accurate analysis of a few lines. We will identify situations for which other models are helpful, e.g., for the case were the emitter-perturber interactions cannot be represented by a sequence of collisions. Such models use the statistical properties of the electric field created by the perturbing particles. In astrophysics, model microfield methods provide an efficient alternative for cases where neither the impact nor the static approximation are valid. For such situations, several models have been developed and interfaced with atomic data. Their accuracy can be tested by simulation techniques avoiding some approximations, but at the expenses of computer time. Such computer simulations can be used to analyze the various physical processes involved in plasmas under arbitrary conditions. We will illustrate their use in the case of plasma rogue waves.

2. Impact Broadening

A detailed and accurate modeling of Stark broadening started almost sixty years ago with the development of a general impact theory having the ability of retaining the quantum character of the emitters and perturbers, and allowing both elastic and inelastic collisions between such particles [2]. The line shape is obtained by a Fourier transform of the dipole autocorrelation function (DAF), a quantity expressed as a trace over all possible states of the quantum emitter plus perturbers system:

$$C(t) = Tr[\vec{d} . T^+(t)\vec{d} T(t)\rho],$$
(1)

where d is the dipole moment of the emitter, $T(t) = exp(-iHt/\hbar)$ the evolution operator and ρ the density matrix, these last two quantities being dependent on the Hamiltonian H for the whole system. Such an expression could be calculated by density functional theory or quantum Monte Carlo methods, taking advantage of the development of computational techniques and computer hardware [3]. Such studies have proved to be efficient for describing the properties of dense plasmas found in the interior of gaseous planets, the atmospheres of white dwarfs or the laboratory plasmas created by energetic lasers. They might be useful for understanding some features in the spectrum of such plasmas, but have not been developed yet in the context of line broadening. Probably the main reason for this is that there is no clear evidence that the dynamical effects of multiple quantum perturbers can affect a line shape. Another reason is that for most of the plasma conditions and line shapes studied, we can use the impact approximation, which assumes that the various perturbers interact separately with the emitter (binary collision assumption), and that the average collision is weak. A validity condition for the impact approximation is that the collision time is small compared with the decorrelation time of C(t). If this condition and the binary collision assumptions are verified, it is possible to use a constant collision operator to account for all the effects of the perturbers on the emitter. Different approaches using impact approximation have been proposed, but we can distinguish firstly between quantum impact models that retain the quantum behavior of both emitters and perturbers, and the semiclassical impact models treating only the emitter as a quantum particle. A pictorial representation of the full quantum emitter-perturber interaction is provided by the use of wave packets for the perturbers. Each wave packet is scattered in a region within the reach of the interaction potential with the emitter. Quantum collision formalism can be applied, enabling the calculation of cross sections with the aid of scattering amplitudes. Although quantum impact calculations have been performed since the seventies [4,5], such calculations are not very numerous for line broadening due to their computational difficulty. In particular, they involve a calculation of the scattering matrix or S matrix [2]. Many calculations have been applied to isolated lines of various ions, a case for which the width w takes the compact form of an average over the perturbers after the use of the optical theorem [2]:

$$w = \frac{1}{2}N\{v[\sigma_i + \sigma_f + \int d\Omega | f_i(\theta, \varphi) - f_f(\theta, \varphi)|^2]\}_{Av},$$
⁽²⁾

where *N* is the perturbers' density, *v* their velocity, σ_i and σ_f are inelastic cross sections, f_i and f_f the forward-scattering amplitudes in a direction given by θ , φ for the initial *i* and final *f* states, and $\{\ldots\}_{Av}$ stands for a Boltzmann thermal average.

With the advent of accurate atomic structure and *S* matrix codes, such impact quantum calculations have been given a new life [6,7], and are most often in good agreement with other calculations. A very efficient calculation has been proposed starting from Equation (2), using a Bethe-Born approximation [8] for evaluating inelastic cross sections. This semi-empirical model uses an effective Gaunt factor, a quantity which measures the probability of an incident electron changing its kinetic energy [9]. This function has been modified and improved to develop the modified semi-empirical model which is frequently used for calculations of isolated ion lines [1].

For most plasma conditions and line shapes studied, the wave packets associated with the perturbers are small and do not spread much in time. This enables the use of classical perturbers following classical paths. Different approaches use this approximation together with the impact approximation for the electron perturbers. Early calculations of hydrogen lines with comparisons to experimental profiles proved that a profile using an impact electron broadening [10], together with a static approximation for the ion perturbers, is in overall agreement for the Balmer H_β line in an arc plasma with a density $N = 2.2 \times 10^{22} \text{ m}^{-3}$ and a temperature T = 10,400 K. The remaining discrepancies concerned the central part of the line and the far line wings, two regions that required an improvement of the model.

For isolated lines of neutral atoms and ions, the semiclassical perturbation (SCP) method [11] has been successfully applied to numerous lines, and is implemented in the STARK-B database [12]. The SCP method was inspired by developments in the quantum theory of collisions between atoms and electrons or ions, and, e.g., performs the angular averages with Clebsch-Gordan coefficients. It has the ability to generate several hundred lines rapidly for a set of densities and temperatures in a single run. The accuracy of the SCP method is assessed by a comparison to experimental spectra, and is about 20 to 30% for the widths of simple spectra, but could be worse for some complex spectra. The method is continuously improving, and has been interfaced with atomic structure codes [11].

An interesting point is raised by the comparison of impact quantum and semiclassical calculations, and a comparison of those with experiments. Quantum calculations have often been found to predict narrower lines than those of semiclassical models [13]. Semiclassical calculations may be brought closer to quantum widths, e.g., by a refined calculation of the minimum impact parameter allowing the use of a classical path [14]. Surprisingly, quantum widths of Li-like and boron-like ions often show a worse agreement with experiments than semiclassical calculation, thus calling for further calculations and analysis [7,15]. As an example of such, more recent quantum calculations [15] are in fairly good agreement with experiments [15].

3. Simulations of Impact Theory and Ion Dynamics

The need for a model that does not assume the impact approximation arose out of the study of hydrogen lines, with the surge of accurate profile measurements in near equilibrium plasmas [16]. It appeared that a standard model using a static ion approximation, and an impact electron collision operator, showed pronounced differences with the measures near the line centers, and also in the far wings. The line wings were well reproduced by the so-called unified theory, which retains the static interaction between an electron and the atom as a strong collision occurs [17,18]. The difference in the central part of the line was linked to the use of the static ion approximation, since it depended on the reduced mass of the emitter-ion perturber system. The observation of the Lyman- α (L $_{\alpha}$) line [19]

showed later that the experimental profile was a factor 2.5 broader than the theoretical line using static ions in arc plasma conditions. This was a strong motivation for developing a technique able to retain ion dynamics in a context where the electric field is created by numerous ions in motion. Since perturbative approaches were unable to account for multiple strong collisions, a computer simulation has been proposed for describing the motion of the ions. The effect on the emitter of the time dependent ion electric field is obtained by a numerical integration of the Schrödinger equation. Early calculations showed the effect of ion dynamics in the central part of the line, and were able to strongly reduce the difference between experimental and simulation profiles [20–22].

Simple hydrogen plasma simulations may be used to illustrate the behavior of an electric field component during a time interval of the order of the line shape time of interest. This time is usually taken as the DAF decorrelation time, and can also be defined as the inverse of the line width. The electric field experienced by an atom surrounded by moving charged particles can be calculated at the center of a cubic box, using particles with straight line trajectories. The edge of the cube should be assumed to be equal to a few times the Debye length $\lambda_D = \sqrt{\epsilon_0 k_B T / (Ne^2)}$, with T and N the hydrogen plasma temperature and density, respectively, e the electron charge, k_B the Boltzmann constant, and ε_0 the permittivity of free space. If we simulate only the ion perturbers, we assume that each particle creates a Debye shielded electric field, in an attempt to retain ion-electron correlations. Random number generators are used to obtain the uniform positions and Maxwell-Boltzmann distributed velocities of the charged particles. If an ion leaves the cubic box, it is replaced by a new one created near the cube boundaries. For the weak coupling conditions assumed, a large number of particles (several thousand commonly) is retained in a cube with a size larger than the Debye length. Such a model provides a good approximation for the time-dependent electric field in a weakly coupled ion plasma at equilibrium, although it suffers from inaccuracies, especially if the size of the box is not large enough [23]. We show in Figure 1 the time dependence of one component of the ionic electric field calculated at the center of the box for an electron density $N_e = 10^{19} \text{ m}^{-3}$, and a temperature T = 40,000 K. The electric field is expressed in units of $E_0 = 1/(4\pi\epsilon_0 r_0^2)$, where r_0 is the average distance between particles defined by $r_0^3 = 3/(4\pi N_e)$. The time interval of 5 ns used in Figure 1a is the L_a time of interest for such plasma conditions. The validity condition of the binary collision approximation requests that the Weisskopf radius $\rho_w = \hbar n^2 / m_e v_i$, with *n* the principal quantum number of the L α upper states (*n* = 2), and $v_i = \sqrt{2k_BT/m_p}$ the thermal ion velocity (m_e and m_p are resp. the electron and proton mass), is much smaller than the average distance between particles. This ratio is for L_{α} and protons of the order of 0.04, enabling the use of an impact approximation. The electric field in Figure 1a clearly exhibits several large fields that are well separated in time during the 5 ns of the L_{α} time of interest. During this time interval, only a few fields (3 in Figure 1a) have a magnitude larger than 50 E_0 , but about 20 have a magnitude of 10 E₀ or more. A piece of the same field history is shown in Figure 1b during a time interval equal to the time of interest for the Balmer- β (H_{β}) line. For this time interval of 0.3 ns, the electric field shows much fewer fluctuations, the atom is no longer submitted to a sequence of sharp collisions, and we can no longer use the impact approximation. This is confirmed by a value of 0.16 for the ρ_w/r_0 ratio, making the use of an impact approximation for this line problematic. Looking now at Figure 1a,b, we can see a background of electric field fluctuations with a small magnitude of about E_0 , and a typical time scale longer than the collision time r_0/v_i . Such fluctuations correspond to the sum of electric fields of distant particles with a magnitude on the order of E_0 . For hydrogen lines affected by the linear Stark effect, it is well known that this effect of weak collisions is dominant in near impact regimes [10], and results from the long range of the Coulomb electric field.



Figure 1. Electric field component in units of E_0 in a plasma with a density $N_e = 10^{19} \text{ m}^{-3}$, and a temperature T = 40,000 K, during (**a**) a time interval of the order of the time of interest for the L_{α} line, and (**b**) a time interval of the order of the time of interest for the H_{β} line.

Using several thousand samples of such electric fields, it is possible to calculate the DAF for each line studied. This requires for each field history $\vec{E}(t)$ an integration of the Schrödinger equation of the emitter submitted to a dipolar interaction potential $-\vec{d} \cdot \vec{E}(t)$. We obtain the emitter's evolution operator by finite difference computational methods, using time steps adjusted to ensure the best compromise between computer time cost and accuracy [24]. The integration time interval is provided by the time of interest for the line calculated, and a first estimate for the time step is a hundredth of the collision time. In the following, we retain only the broadening of the upper states of the line, resulting in some loss of accuracy for the first Balmer lines, but in a much faster calculation. We show in Figure 2a the DAF of L_{α} for the same plasma conditions as in Figure 1. The ab-initio DAF (solid line) is obtained by a simulation of the ions retaining also the effect of electrons with an impact approximation. We observe that this simulation is close to an impact calculation for both ions and electrons (dashed line). For the same condition and the H_{β} line, the decay of the ab-initio DAF is significantly smaller than for the impact calculation, indicating again a deviation from ion impact broadening for this line.



Figure 2. Dipole autocorrelation functions with an ab-initio simulation (solid line) and in the impact limit (dotted line) in a plasma with a density $N_e = 10^{19} \text{ m}^{-3}$, and a temperature T = 40,000 K, for (**a**) the L_{α} Lyman transition, and (**b**) the H_{β} Balmer transition.

Another way of taking account of ion dynamics is with the help of stochastic processes. A stepwise constant stochastic process is used to model the electric field felt by the atom [25]. The process requires the knowledge of the microfield probability distribution function, and of a waiting time distribution function controlling the jumps from one field to the next one. Such model microfield methods are efficient for retaining ion dynamics effects, and are used for a diagnostic of hydrogen lines [26]. Stochastic processes are also used in the line shape code using the frequency fluctuation model for an inclusion of ion dynamics [27]. During the last decades, several simulations and models have been developed with the ability of retaining ion dynamics. The field is still active, with ion dynamics being one of the issues discussed in the Spectral Line Shapes in Plasmas workshop, providing many new analyses [28].

4. Effect of Plasma Waves

Plasmas sustain various types of waves, which behave differently in a linear and nonlinear regime. A way to distinguish between the two regimes is to calculate the ratio W of the wave energy density to the plasma energy density, given by:

$$W = \varepsilon_0 E_L^2 / 4N_e k_B T , \qquad (3)$$

where E_L is the electric field magnitude of the wave. For values of *W* much smaller than 1, we expect a linear behavior of the waves. In a linear regime, electronic Langmuir waves oscillate at a frequency close to the plasma frequency $\omega_p = \sqrt{N_e e^2/m_e \varepsilon_0}$, and can be excited even by thermal fluctuations. We assume that the numerous emitters on the line of sight are submitted to different Langmuir waves, each with the same frequency ω_p , but a different direction and phase chosen at random, and a magnitude sampled using a half-normal probability density function (PDF). In the following, we have used this half-normal PDF for the reduced electric field magnitude $F = E/E_0$:

$$P(F) = \frac{\sqrt{2}}{\sigma\sqrt{\pi}} exp\left(-\frac{F^2}{2\sigma^2}\right) \tag{4}$$

In this expression, we use the standard deviation σ of a normal distribution, and thus obtain the mean value E_L of E by writing $E_L = \sigma E_0 \sqrt{2/\pi}$. Each Langmuir wave has a different electric field history, and we obtain the DAF by an average over about a thousand such field histories. For a plasma with a density $N_e = 10^{19} \text{ m}^{-3}$, and a temperature $T = 10^5 \text{ K}$, we first calculated the L_{α} DAF for Langmuir waves with a mean electric field magnitude corresponding to W = 0.01 ($E_L = 15E_0$). The response of the DAF is a periodic oscillation with a period equal to $2\pi/\omega_p$, but with an amplitude much smaller than 1 for this average field magnitude of $15E_0$. After a product with an impact DAF for retaining the effect of the background electron and ion plasma, there remains no visible effect of the waves on the convolution DAF for the value W = 0.01. This ratio can take much larger values, however; especially if an external energy source such as a beam of charged particles is present. As W increases, nonlinear phenomena start showing up, enabling, for instance, wave-wave couplings. Although only recently investigated in plasmas, the occurrence of rogue waves has been raised in various plasma conditions [29–31]. Rogue waves have been studied in many dynamical systems, and are known to the general public by the observation and study of rogue or freak waves that suddenly appear in the ocean as large isolated waves. In oceanography, rogue waves are defined as waves whose height is more than twice the mean of the largest third of the waves in a wave record. Rogue waves appear to be a unifying concept for studying localized excitations that exceed the strength of their background structures. They are studied in nonlinear optics [32], Bose-Einstein condensates [33], and many other fields outside of physics. For our line shape problem in plasmas, we postulate that nonlinear processes create rogue waves from a random background of smaller Langmuir waves. The physical mechanism at play is the coupling of the Langmuir wave with ion sound and electromagnetic waves; density fluctuations of the sound waves affect the high frequency waves through ω_p . The first Zakharov equation [34] shows how

density fluctuations affect Langmuir waves, and a second equation how a Langmuir wave packet can produce a density depression via the ponderomotive force [35]. We will not discuss these equations here, which are particularly useful for a study of wave collapse. Most present rogue wave studies rely on the nonlinear Schrödinger equation (NLSE), which is obtained in the adiabatic limit (slowly changing density perturbations) of the Zakharov equations [35]. A one-dimensional solution of the NLSE is commonly used to approximate the response of nonlinear media. Stable envelope solitons are possible solutions of the 1D NLSE. We will assume that there is a contribution of a stable envelope soliton for each history of the microfield, similarly to what we did for the background Langmuir wave. Using a ratio W = 0.1, the average peak magnitude of such solitons will be 3 times the amplitude of background Langmuir waves, fitting them in the category of rogue waves. A possible shape for the envelope is a Lorentzian, with a time dependence that bears some similarity with the celebrated Peregrine soliton [36]. We observe in Figure 3 that the DAF of L_{α} obtained with a product of the impact DAF and the Langmuir rogue wave DAF for W = 0.1 is affected by oscillations at the plasma frequency.



Figure 3. L_{α} dipole autocorrelation function in a plasma with a density $N_e = 10^{19} \text{ m}^{-3}$, and a temperature $T = 10^5$ K, calculated with a product of the impact DAF and the Langmuir rogue wave DAF for W = 0.1.

Looking at the line shape obtained with a Fourier transform, we can see in Figure 4 that the peak of the line including the wave effect is about 10% lower than the impact line peak, but this with almost no effect on the line width. Not shown in Figure 4, we noticed that the wing of the line affected by rogue waves had a slightly slower decay than the impact profile, indicating a transfer of intensity from the center toward large line shifts. It is remarkable that a such rogue wave had a rather small effect on the profile. This is probably due to the fact that we are in impact conditions for this line. In impact regimes, decorrelation is very effective, leaving only a small broadening contribution to the type of rogue waves that we considered. A larger broadening effect would be observed by considering wave collapse, a phenomenon occurring as *W* takes larger values of the order of 1 or more for such plasma conditions. The emitters then experience the effect of a sequence of solitons which can significantly increase the broadening [37].



Figure 4. L_{α} in a plasma with a density $N_e = 10^{19} \text{ m}^{-3}$, and a temperature $T = 10^5$ K, calculated with an impact approximation (dashed line), and with a Fourier transform of the DAF in Figure 3 (solid line).

5. Conclusions

Impact approximation mainly consists of saying that, on an average, it takes many collisions to change the quantum state of an atom. When this approximation is valid, the effect of the numerous fluctuating interactions of the emitter with the perturbers can be expressed with a constant collision operator. We have briefly described several models using impact approximation. A wide variety of impact models have been proposed, ranging from full quantum calculations to semiclassical approaches. Impact calculations allow expression of the width and shift of a line in terms of quantum scattering cross-sections. Such calculations have enabled many improvements in the application of quantum theory for obtaining observable quantities such as a line shape. The comparison between experimental and theoretical spectra is of great benefit for the validation of such models. It is thus crucial to be able to rapidly obtain numerous spectra for the lines of many atoms and ions. This is possible using models such as the semiclassical perturbation or the semi-empirical formalism. We have also shown how a computer simulation can reproduce the results of the impact approximation for hydrogen lines. Such simulations involve several thousand particles, however, and are certainly not the most efficient technique for obtaining the impact profile. The main advantage of simulations is that they can go beyond the impact approximation, for situations with many perturbers acting simultaneously on the emitter. We have briefly recalled the problem of ion dynamics, and have proposed a simple simulation for a calculation of the effect of Langmuir rogue waves of L_{α} in an impact regime.

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Article Quantum and Semiclassical Stark Widths for Ar VII Spectral Lines

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Abstract: We present in this paper the results of a theoretical study of electron impact broadening for several lines of the Ar VII ion. The results have been obtained using our quantum mechanical method and the semiclassical perturbation one. Results are presented for electron density 10^{18} cm⁻³ and for electron temperatures ranging from 2×10^4 to 5×10^5 K required for plasma modeling. Our results have been compared to other semiclassical ones obtained using different sources of atomic data. A study of the strong collisions contributions to line broadening has been performed. The atomic structure and collision data used for the calculations of line broadening are also calculated by our codes and compared to available theoretical results. The agreement found between the two calculations ensures that our line broadening procedure uses adequate structure and collision data.

Keywords: Stark broadening; Ar VII line profiles; stars; white dwarfs; atomic data; scattering

1. Introduction

Atomic and line broadening data for many elements and their ions are very useful for solving many astrophysical problems, such as the calculations of opacity and radiative transfer [1]. Especially, accurate Stark broadening parameters are important to obtain a reliable modelization of stellar interiors. The Stark broadening mechanism is also important for the investigation, analysis, and modeling of B-type and particularly A-type stellar atmospheres, as well as white dwarf atmospheres [2,3]. Furthermore, the development of computers and instruments, such as the new X-ray space telescope *Chandra*, has motivated the calculations of line broadening of trace elements in the X-ray wavelength range. It has been shown that analysis of white dwarf atmospheres, where Stark broadening is dominant compared to the thermal Doppler broadening, needs models taking into account heavy element opacity.

In Rauch et al. [4], the authors reported problems encountered in their determination of element abundances: the line cores of the S VI resonance doublet appear too deep to match the observation and they are not well suited for an abundance determination, and the same problem exists in relation to the N V and O VI. This is due to the lack of line broadening data for these ions. Some other data exist, but the required temperatures and electron densities are lacking, and it is necessary to extrapolate such data to obtain the temperatures and densities at the line-forming regions, especially the line cores. This procedure of extrapolation can provide inaccurate results especially in the case of extrapolating to obtain temperatures, since the temperature dependence of line widths may be very different. This lack of data represents an inconvenience for the development of spectral analysis by means of the NLTE

model atmosphere techniques. We quote here the conclusion in Rauch et al. [4]: "spectral analysis by means of NLTE model atmospheres has presently arrived at a high level of sophistication, which is now hampered largely by the lack of reliable atomic data and accurate line-broadening tables."

Astrophysical interest of Ar VII illustrates for example recent discovery of far UV lines of this ion in the spectra of very hot central stars of planetary nebulae and white dwarfs [5]. In this article, the authors have also shown the importance of the line broadening data for this element in its various ionization stages. Argon also has an important role in plasma technological applications and devices [6]. It produces favorable conditions for very stable discharges and is also very often used as a carrier gas in plasma, which contains a mixture of other gases. Thus, the knowledge of the Stark broadening parameters of neutral and ionized argon lines is an important tool for plasma electron density diagnostic.

The Stark broadening calculations in the present work are based on two approaches: the quantum mechanical approach and the semiclassical perturbation one. The quantum mechanical expression for electron impact broadening calculations for intermediate coupling was obtained in Elabidi et al. [7]. The first applications were performed for the 2s3s–2s3p transitions in Be-like ions from nitrogen to neon [8] and for the 3s–3p transitions in Li-like ions from carbon to phosphor [9]. This approach was also used in Elabidi & Sahal-Bréchot [10] to check the dependence on the upper level ionization potential of electron impact widths and in Elabidi et al. [11] to investigate the influence of strong collisions and quadrupolar potential contributions on line broadening. Our quantum approach is an ab initio method; i.e., all the parameters required for the calculations of the line broadening such as radiative atomic data (energy levels, oscillator strengths ...) or collisional data (collision strengths or cross sections, scattering matrices ...) are evaluated during the calculation and not taken from other data sources. We used the sequence of the University College London (UCL) atomic codes SUPERSTRUCTURE/DW/JAJOM that have been used for many years to provide fine energy levels, wavelengths, radiative probability rates, and electron impact collision strengths. Recently, they have been adapted to line broadening calculations [8].

In the present paper, we continue the effort to provide atomic and line broadening data for argon ions. Quantum Stark broadening of 12 lines of the Ar VII ion have been calculated using 9 configurations $(1s^22s^22p^6: 3s^2, 3s3p, 3p^2, 3s3d, 3p3d, 3s4s, 3s4p, 3s4d, and 3s5s)$. Our calculations have been made for a set of temperatures ranging from 2×10^4 to 5×10^5 K. These parameters will be useful for a more accurate determination of photospheric properties. We perform also a semiclassical calculations for these lines using our atomic data from the code SUPERSTRUCTURE. We compare these results to the semiclassical ones [12], for which the atomic structure has been calculated with the Bates and Damgaard approximation [13].

2. Outline of the Theory and Computational Procedure

2.1. Quantum Mechanical Formalism

We present here an outline of our quantum formalism for electron impact broadening. More details can be found elsewhere [7,8]. The calculations are made within the frame of the impact approximation, which means that the time interval between collisions is much longer than the duration of a collision. The expression of the Full Width at Half Maximum (FWHM) W obtained in Elabidi et al. [8] is:

$$W = 2N_e \left(\frac{\hbar}{m}\right)^2 \left(\frac{2m\pi}{k_BT}\right)^{\frac{1}{2}} \times \int_{0}^{\infty} \Gamma_w(\varepsilon) \exp\left(-\frac{\varepsilon}{k_BT}\right) d\left(\frac{\varepsilon}{k_BT}\right)$$
(1)

where k_B is the Boltzmann constant, N_e the electron density, T the electron temperature, and

$$\Gamma_{w}(\varepsilon) = \sum_{J_{i}^{T} J_{f}^{T} l K_{i} K_{f}} \frac{\left[K_{i}, K_{f}, J_{i}^{T}, J_{f}^{T}\right]}{2} \\ \times \left\{ \begin{array}{c} J_{i} K_{i} l \\ K_{f} J_{f} 1 \end{array} \right\}^{2} \left\{ \begin{array}{c} K_{i} J_{i}^{T} s \\ J_{f}^{T} K_{f} 1 \end{array} \right\}^{2} \\ \times \left[1 - (\operatorname{Re}\left(S_{I}\right) \operatorname{Re}\left(S_{F}\right) + \operatorname{Im}\left(S_{I}\right) \operatorname{Im}\left(S_{F}\right))\right] \end{array}$$
(2)

where $L_i + S_i = J_i$, $J_i + l = K_i$ and $K_i + s = J_i^T$. L and S represent the atomic orbital angular momentum and spin of the target, *l* is the electron orbital momentum, and the superscript *T* denotes the quantum numbers of the total electron+ion system. $S_I(S_F)$ are the scattering matrix elements for the initial (final) levels, expressed in the intermediate coupling approximation, Re (S) and Im (S) are respectively the

real and the imaginary parts of the S-matrix element, $\begin{cases} abc \\ def \end{cases}$ represent 6–j symbols, and we adopt the notation [x, y] = (2x + 1)(2y + 1). Both S_I and S_F are calculated for the same incident electron energy $\varepsilon = mv^2/2$. Equation (1) takes into account the fine structure effects and relativistic corrections resulting from the breakdown of the *LS* coupling approximation for the target.

The main goal is the evaluation of the real (Re **S**) and the imaginary parts (Im **S**) of the scattering matrix **S** in the initial *I* and the final *F* level. The calculation starts with the study of the atomic structure. The structure problem has been treated using the SUPERSTRUCTURE (SST) code described in Eissner et al. [14], taking into account configuration interaction, where each individual configuration is an expansion in terms of Slater states built from orthonormal orbitals. The radial functions were calculated assuming a scaled Thomas–Fermi–Dirac–Amaldi (TFDA) potential. The potential depends upon parameters λ_I which are determined variationally by optimizing the weighted sum of the term energies. Relativistic corrections (spin-orbit, mass, Darwin, and one-body) are introduced according to the Breit–Pauli approach [15] as a perturbation to the non-relativistic Hamiltonian. The SST program also produces the term coupling coefficients (TCCs), which are used to transform the scattering **S** or reactance **R**-matrices to intermediate coupling [7].

The second step is the treatment of the scattering problem. The calculation is carried out in the non-relativistic distorted wave approximation using the UCL distorted wave (DW) program [16]. The reactance matrices are calculated in LS coupling. The program JAJOM [17] uses these reactance matrices and the TCC to calculate collision strengths in intermediate coupling. In the present work, we have transformed JAJOM into JAJPOLARI (Elabidi and Dubau, unpublished results) to produce the collision strengths and the reactance matrices **R** in intermediate coupling, which will be used by the program RtoS (Dubau, unpublished results) to evaluate the real and the imaginary parts of the scattering matrix according to

$$\operatorname{Re} \mathbf{S} = \left(1 - \mathbf{R}^2\right) \left(1 + \mathbf{R}^2\right)^{-1}$$
(3)

and

$$\operatorname{Im} \mathbf{S} = 2\mathbf{R} \left(1 + \mathbf{R}^2 \right)^{-1}.$$
 (4)

The two expressions (3) and (4) have been deduced from the relation $S = (1 + iR)(1 - iR)^{-1}$, and such expressions guarantee the unitarity of the S-matrix.

Finally, in the code JAJPOLARI, the reactance matrices R_I in intermediate coupling corresponding to the initial I level are evaluated for each channel and at a total energy $E_I = E_i + \varepsilon$. The same procedure is done for R_F but at a total energy $E_F = E_f + \varepsilon$. E_i (E_f) are the energies of the initial (final) atomic levels. The program RtoS receives **R**-matrices and transforms them into real and imaginary parts of **S**-matrices according to Equations (3) and (4) at total energies E_I and E_F , and combines a given matrix element S_I for an initial level I with a number of matrix element S_F for the final level F. The obtained matrix elements Re **S** and Im **S** enter into Equation (2). The integral over the Maxwell distribution (Equation (1)) is evaluated numerically using a trapezoid integration with a variable step to provide the line width *W*. The energy step is chosen to be as small as possible around the threshold region where the variation of Γ_w in (1) is fast. For large energies and far from the threshold region, the variation of Γ_w becomes slow and then the step is gradually increased.

2.2. Semiclassical Perturbation Method

We give here a detailed description of the semiclassical perturbation formalism for line broadening calculations. The profile $F(\omega)$ is Lorentzian for isolated lines:

$$F(\omega) = \frac{w/\pi}{(\omega - \omega_{if} - d)^2 + w^2}$$
(5)

where

$$\omega_{if} = \frac{E_i - E_f}{\hbar}$$

i and *f* denote the initial and final atomic states and E_i and E_f their corresponding energies.

The total width at half maximum (W = 2w) in angular frequency units of a spectral line can be expressed as

$$W = N \int v f(v) dv \left(\sum_{i' \neq i} \sigma_{ii'}(v) + \sum_{f' \neq f} \sigma_{ff'}(v) + \sigma_{el} \right)$$
(6)

where *N* is the electron density, f(v) the Maxwellian velocity distribution function for electrons, i' (resp. f') denotes the perturbing levels of the initial state *i* (resp. final state *f*). The inelastic cross section $\sigma_{ii'}(v)$ (resp. $\sigma_{ff'}(v)$) can be expressed by an integral over the impact parameter ρ of the transition probability $P_{ii'}(\rho, v)$ (resp. $P_{ff'}(\rho, v)$) as

$$\sum_{i'\neq i} \sigma_{ii'}(v) = \frac{1}{2}\pi R_1^2 + \int_{R_1}^{R_D} 2\pi\rho d\rho \sum_{i'\neq i} P_{ii'}(\rho, v)$$
(7)

where ρ denotes the impact parameter of the incoming electron. The elastic cross section is given by

$$\sigma_{el} = 2\pi R_2^2 + \int_{R_2}^{R_D} 2\pi\rho d\rho \sin^2 \delta + \sigma_r$$

$$\delta = (\varphi_p^2 + \varphi_q^2)^{\frac{1}{2}}.$$
(8)

Strong collisions are evaluated for $\rho < R_1, R_2$. The phase shifts φ_p and φ_q , due respectively to the polarization potential (r^{-4}) and to the quadrupolar potential (r^{-3}), are given in Section 3 of Chapter 2 in Sahal-Bréchot [18], and R_D is the Debye radius. The cut-offs R_1 and R_2 are described in Section 1 of Chapter 3 in Sahal-Bréchot [19]. Detailed calculations of the interference term σ_{el} can be found in Formulas 18 and 24–30 on pages 109–110 of Sahal-Bréchot [18]. σ_r is the contribution of the Feshbach resonances [20], which concerns only ionized radiating atoms colliding with electrons. It is an extrapolation of the excitation collision strengths (and not the cross-sections) under the threshold by means of the semiclassical limit of the Gailitis approximation (see page 601 of [20] for details of the calculations). A review of the theory, all approximations and the details of applications are given in Sahal-Bréchot et al. [21].

3. Results and Discussions

3.1. Atomic Structure and Electron Scattering Data

We have used the following nine configurations in our calculation: $1s^22s^22p^6$: $3s^2$, 3s3p, $3p^2$, 3s3d, 3p3d, 3s4s, 3s4p, 3s4d, and 3s5s, which give rise to 38 levels, which are listed in Table 1 with their energies in cm⁻¹. These values have been compared with the observed ones taken from the tables of the National Institute of Standards and Technology database: NIST [22] which are originally from Saloman [23]. We compare also with the energies computed using the multiconfiguration Hartree–Fock method (MCHF) [24] and with those obtained using the AUTOSTRUCTURE code [25]. The averaged disagreement between these three results is less than 1%. We detect an inversion between the two levels 10/13 and 25/26 regarding those of NIST and MCHF. This inversion does not affect the calculations since the agreement is still acceptable (about 5%).

Table 1. Our present fine-structure energy levels E (in cm^{-1}) for Ar VII compared with those of NIST [22], with those obtained from the multiconfiguration Hartree–Fock method (MCHF) [24], and with those from the R-matrix calculation (AS2014) [25]. Levels denoted by asterisks (*) are inverted compared to the NIST values.

i	Conf.	Level	Е	NIST	MCHF	AS2014	$\frac{ E-NIST }{NIST}$ (%)
1	$3s^{2}$	${}^{1}S_{0}$	0.0	0.0	0.0	0.0	_
2	3s3p	$^{3}P_{0}^{0}$	110,717	113,101	112,817.66	112,070	2.1
3	3s3p	${}^{3}P_{1}^{0}$	111,488	113,906	113,632.14	112,889	2.1
4	3s3p	$^{3}P_{2}^{0}$	113,088	115,590	115,324.84	114,593	2.2
5	3s3p	${}^{1}P_{1}$	172,878	170,722	170,598.08	173,751	1.3
6	$3p^{2}$	$^{1}D_{2}$	263,439	264,749	264,797.88	264,530	0.5
7	$3p^{2}$	$^{3}P_{0}^{0}$	271,494	269,836	269,688.15	270,704	0.6
8	$3p^{2}$	${}^{3}P_{1}^{0}$	272,341	270,777	270,667.14	271,641	0.6
9	$3p^{2}$	$^{3}P_{2}^{0}$	273,971	272,562	272,474.76	273,432	0.5
10	3s3d	${}^{3}D_{1}^{2}$	325,254	324,104	324,950.35	326,054	0.4
11	3s3d	$^{3}D_{2}$	325,335	324,141	324,966.00	326,141	0.4
12	3s3d	³ D ₃	325,456	324,205	325,056.68	326,273	0.4
13	$3p^{2}$	${}^{1}S_{0}$	333,116	316,717	317,014.73	320,974 *	5.2
14	3s3d	$^{1}D_{2}$	384,031	370,294	371,275.29	377,167	3.7
15	3p3d	${}^{3}F_{2}^{0}$	443,952	443,362	444,508.36	444,677	0.1
16	3p3d	³ F ₃₀	444,892	444,780	445,556.29	445,701	0.0
17	3p3d	${}^{3}F_{4}^{0}$	446,051	446,011	446,849.87	446,969	0.0
18	3p3d	${}^{1}D_{2}$	450,025	450,477	450,808.06	451352	0.1
19	3p3d	$^{3}P_{2}^{0}$	474,314	472,282	473,009.27	475,022	0.4
20	3p3d	${}^{3}P_{1}^{5}$	474,956	472,875	473,782.67	475,699	0.4
21	3p3d	$^{3}P_{0}^{0}$	475,497	473,810	474,466.36	476,301	0.4
22	3p3d	${}^{3}D_{1}$	477,133	475,217	475,932.22	477,901	0.4
23	3p3d	³ D ₂	477,515	475,585	476,306.50	478,313	0.4
24	3p3d	³ D ₃	477,753	475,762	476,474.91	478,560	0.4
25	3s4s	${}^{3}S_{1}$	513,685	514,076	508,971.69	511,372	0.1
26	3p3d	$^{1}F_{3}^{0}$	521,897	510,268	514,890.47	515,169 *	2.3
27	3p3d	${}^{1}P_{1}^{0}$	527,518	517,105	517,788.24	524,282	2.0
28	3s4s	${}^{1}S_{0}$	529,866	528,910	526,205.45	523,618	0.2
29	3s4p	$^{3}P_{0}^{0}$	567,050	563,880	568,040.66	565,087	0.6
30	3s4p	$^{3}P_{1}^{0}$	567,287	564,418	568,275.74	565,295	0.5
31	3s4p	$^{3}P_{2}^{0}$	567,811	564,728	568,944.94	565,840	0.5
32	3s4p	${}^{1}P_{1}^{\bar{0}}$	576,576	569,797	570,403.78	568,205	0.2
33	3s4d	${}^{3}D_{1}$	635,209	634,605	635,580.25	632,497	0.1
34	3s4d	³ D ₂	635,241	634,639	635,659.10	632,562	0.1
35	3s4d	³ D ₃	635,290	634,701	635,749.02	632,659	0.1
36	3s4d	${}^{1}D_{2}$	639,087	635,295	636,353.38	633,443	0.6
37	3s5s	${}^{3}S_{1}$	713,912	715,747	_	717,638	0.3
38	3s5s	${}^{1}S_{0}$	719,473	714,794	_	717,997	0.7

We present also in Table 2 radiative decay rates A_{ij} , weighted oscillator strengths gf, and line strengths S for some Ar VII lines up to the level 14 (3s3d ¹D₂). Our A_{ij} values have been compared with those obtained from the AUTOSTRUCTURE code [25], and with those from the SUPERSTRUCTURE code [26] using five configurations ($1s^22s^22p^6$: $3s^2$, 3s3p, $3p^2$, 3s3d, and 3s4s). The averaged difference is about 20% with the results of [25] and about 24% with those of Christensen et al. [26]. Some transitions present a high difference, especially those for which A_{i-j} are relatively small (about 10^6 s^{-1} and below). The gf values have been compared only with Christensen et al. [26] and the difference is about 24 %. The gf values are calculated in [26], but we took them from the database CHIANTI version 8.0 [27].

With the code JAJOM, fine structure collision strengths are calculated for low partial waves l of the incoming electron up to 29. For large partial waves l, this method becomes cumbersome and inaccurate, but their contributions to collision strengths cannot be neglected. For $30 \le l \le 50$, two different procedures have been used: for dipole transitions, the contribution has been calculated using the JAJOM-CBe code (Dubau, unpublished results) based upon the Coulomb–Bethe formulation of Burgess and Sheorey [28] and adapted to JAJOM approximation. For non-dipole transitions, the contribution has been estimated by the SERIE-GEOM code assuming a geometric series behavior for high partial wave collision strengths [29,30].

Table 2. Present radiative decay rates A_{ij} (in s⁻¹) and weighted oscillator strengths gf compared to those from Christensen et al. [26] (SST86) and to those from [25] (AS2014) for some Ar VII allowed transitions. Line strengths *S* are also presented. *i* and *j* label the levels as in Table 1.

i - j	A_{i-j}	$A_{i-j}(AS2014)$	$A_{i-j}($ SST86 $)$	gf	gf(SST86)	S
3 - 1	$5.968 imes 10^5$	7.13×10^5	1.65×10^5	$2.160 imes 10^{-4}$	$5.820 imes 10^{-5}$	0.000638
5 - 1	8.114×10^9	8.30×10^9	8.21×10^9	1.221×10^{0}	1.270×10^{0}	2.325423
6 - 3	1.854×10^7	2.97×10^{7}	2.39×10^{7}	$6.018 imes10^{-3}$	$7.780 imes 10^{-3}$	0.013038
6 - 4	3.748×10^7	$6.18 imes 10^7$	$6.07 imes 10^7$	$1.243 imes10^{-2}$	$2.020 imes 10^{-2}$	0.027216
6 - 5	$3.719 imes 10^8$	$4.00 imes 10^8$	$3.98 imes 10^8$	$3.400 imes10^{-1}$	3.380×10^{-1}	1.235825
7 - 3	7.249×10^9	$6.94 imes 10^9$	$6.93 imes 10^9$	$4.245 imes10^{-1}$	$4.280 imes10^{-1}$	0.873472
7 - 5	7.818×10^5	1.72×10^{6}	1.02×10^{6}	1.205×10^{-4}	1.590×10^{-4}	0.000402
8 - 2	2.492×10^{9}	$2.40 imes 10^9$	$2.35 imes 10^9$	4.291×10^{-1}	4.250×10^{-1}	0.874104
8 - 3	1.842×10^9	1.77×10^9	1.74×10^9	$3.202 imes 10^{-1}$	$3.180 imes10^{-1}$	0.655396
8 - 4	2.976×10^9	2.85×10^9	$2.84 imes 10^9$	$5.278 imes 10^{-1}$	$5.310 imes10^{-1}$	1.090986
8 - 5	1.274×10^5	$1.33 imes 10^5$	$3.65 imes 10^4$	5.793×10^{-5}	1.670×10^{-5}	0.000192
9 - 3	1.877×10^{9}	1.80×10^9	1.77×10^9	5.329×10^{-1}	5.270×10^{-1}	1.079810
9 - 4	5.481×10^9	5.24×10^9	5.22×10^9	1.587×10^{0}	1.590×10^{0}	3.248175
9 - 5	$4.965 imes 10^6$	$8.35 imes 10^6$	7.11×10^{6}	3.642×10^{-3}	5.240×10^{-3}	0.011859
10 - 2	5.993×10^{9}	5.86×10^{9}	5.92×10^{9}	5.857×10^{-1}	5.980×10^{-1}	0.898754
10 - 3	4.449×10^{9}	4.35×10^9	$4.40 imes 10^9$	4.379×10^{-1}	4.480×10^{-1}	0.674448
10 - 4	2.906×10^{8}	$2.84 imes 10^8$	$2.88 imes 10^8$	2.904×10^{-2}	2.980×10^{-2}	0.045058
10 - 5	5.117×10^{5}	5.88×10^5	1.52×10^{5}	9.913×10^{-5}	2.950×10^{-5}	0.000214
11 - 3	$8.015 imes 10^9$	7.84×10^9	7.92×10^9	1.314×10^{0}	1.340×10^{0}	2.022700
11 - 4	2.618×10^{9}	2.56×10^{9}	$2.60 imes 10^9$	4.356×10^{-1}	4.480×10^{-1}	0.675718
11 - 5	7.418×10^5	7.91×10^5	1.15×10^5	$2.392 imes 10^{-4}$	$3.710 imes 10^{-5}$	0.000517
12 - 4	$1.048 imes 10^{10}$	1.02×10^{10}	$1.04 imes 10^{10}$	2.439×10^{0}	2.510×10^{0}	3.781547
13 - 3	4.347×10^{6}	9.07×10^{6}	5.79×10^{6}	1.327×10^{-4}	$2.030 imes 10^{-4}$	0.000197
13 - 5	8.643×10^{9}	6.98×10^{9}	6.97×10^9	$5.047 imes 10^{-1}$	$4.840 imes10^{-1}$	1.036879
14 - 3	9.459×10^{6}	$9.90 imes 10^6$	1.40×10^{6}	9.546×10^{-4}	1.590×10^{-4}	0.001153
14 - 4	4.427×10^5	4.77×10^5	4.47×10^5	4.521×10^{-5}	$5.130 imes 10^{-5}$	0.000055
14 - 5	2.085×10^{10}	1.90×10^{10}	$1.88 imes 10^{10}$	3.506×10^{0}	3.540×10^{0}	5.466647

We present our collision strengths from the lowest five levels to the first 14 levels in Table 3 at electron energy values 7.779, 13.674, and 23.336 Ry. We compared them with the 5-configurations collision strengths of Christensen et al. [26]. Some important discrepancies exist for transitions involving levels arising from the 3p³ configuration (levels 7, 8, and 9). Except for these transitions, the agreement (averaged over the three energies and all the other transitions) is about 20%. The agreement between our results and those of [26] is the worse for the electron energy 7.779 Ry. This energy is close to the excitation energy of the last calculated level (here the energy 6.80 Ry of the level 38). In this situation, the contribution of

elastic collisions (which are mostly due to close/strong collisions) is important. We remark also that the agreement is better for transitions from higher levels: for example, $\Delta\Omega_{ij}$ is about 39% for transitions from the level i = 1, and it is about 15% for transitions from the levels i = 4, 5. We note that, in [26], calculations have been carried out for partial waves $l \leq 11$. This may be the origin of the above disagreement for some transitions (we have taken into account partial waves up to 50 in the present work). The difference in the configurations number may also affect the collision strength values.

Table 3. Our collision strengths Ω_{ij} (Present) and those from [26] (DW86) where $1 \le i \le 5$ and $i + 1 \le j \le 14$. *i* and *j* label the levels as in Table 1.

	7.779	Ry	13.67	4 Ry	23.336 Ry	
1 – J	Present	DW86	Present	DW86	Present	DW86
1 - 2	8.901×10^{-3}	$9.29 imes 10^{-3}$	4.523×10^{-3}	$4.54 imes10^{-3}$	$2.067 imes 10^{-3}$	$2.01 imes 10^{-3}$
1 - 3	2.936×10^{-2}	2.87×10^{-2}	1.652×10^{-2}	1.46×10^{-2}	9.202×10^{-3}	7.08×10^{-3}
1 - 4	4.430×10^{-2}	4.64×10^{-2}	2.251×10^{-2}	2.26×10^{-2}	1.029×10^{-2}	1.00×10^{-2}
1 - 5	8.841×10^{0}	8.60×10^{0}	9.848×10^{0}	1.03×10^{1}	1.006×10^{1}	1.21×10^{1}
1 - 6	4.208×10^{-1}	3.57×10^{-1}	4.712×10^{-1}	3.48×10^{-1}	5.067×10^{-1}	3.17×10^{-1}
1 - 7	5.400×10^{-5}	1.06×10^{-4}	2.900×10^{-5}	5.40×10^{-5}	1.200×10^{-5}	2.33×10^{-5}
1 - 8	1.560×10^{-4}	2.81×10^{-4}	8.400×10^{-3}	1.30×10^{-4}	3.300×10^{-3}	4.64×10^{-3}
1-9	4.102×10^{-3}	5.74×10^{-3}	4.449×10^{-3}	5.38×10^{-3}	4.785×10^{-3}	4.80×10^{-3}
1 - 10	2.239×10^{-2}	2.39×10^{-2}	1.051×10^{-2}	1.10×10^{-2}	4.603×10^{-3}	4.73×10^{-3}
1 - 11	3.731×10^{-2}	3.99×10^{-2}	1.752×10^{-2}	1.84×10^{-2}	7.671×10^{-3}	7.87×10^{-3}
1 - 12	5.222×10^{-2}	5.58×10^{-2}	2.452×10^{-2}	2.57×10^{-2}	1.074×10^{-2}	1.10×10^{-2}
1 - 13	5.460× 10 ⁻¹	1.28×10^{-1}	1.030×10 ⁻¹	1.10×10^{-1}	5.800×10^{-1}	8.06×10^{-1}
1 - 14	5.431×10^{-2}	7.40×10^{-2}	6.408×10^{-2}	8.03×10^{-2}	7.164×10^{-2}	8.46×10^{-2}
2 - 3 2 4	0.237×10^{-1}	9.55×10^{-1}	3.113×10^{-1}	3.26×10^{-1}	1.475×10^{-1}	3.05×10^{-1}
2 - 4 2 - 5	1.103×10^{-2}	2.98×10^{-2} 1.19×10^{-2}	5.302×10^{-3}	2.79×10^{-3}	3.333×10^{-3}	2.30×10^{-3}
2 - 5 2 - 6	2.860×10^{-2}	1.17×10^{-2} 2.77×10^{-2}	1.381×10^{-2}	1.40×10^{-2}	5.991×10^{-3}	6.05×10^{-3}
2 - 7	5.657×10^{-3}	2.09×10^{-2}	2.784×10^{-3}	2.95×10^{-3}	1.210×10^{-3}	1.27×10^{-3}
2 - 8	3.379×10^{0}	3.32×10^{0}	3.796×10^{0}	4.05×10^{0}	3.867×10^{0}	4.70×10^{0}
2 - 9	4.773×10^{-3}	6.85×10^{-2}	2.338×10^{-3}	6.45×10^{-3}	1.012×10^{-3}	3.57×10^{-3}
2 - 10	2.744×10^{0}	2.75×10^{0}	3.272×10^{0}	3.43×10^{0}	3.563×10^{0}	4.10×10^{0}
2 - 11	2.286×10^{-2}	4.89×10^{-2}	1.027×10^{-2}	1.23×10^{-2}	4.178×10^{-3}	$5.19 imes 10^{-3}$
2 - 12	$5.517 imes 10^{-2}$	$6.10 imes 10^{-2}$	5.424×10^{-2}	$5.58 imes 10^{-2}$	$5.713 imes 10^{-2}$	$5.14 imes 10^{-2}$
2 - 13	$1.746 imes 10^{-3}$	$1.68 imes 10^{-3}$	$8.200 imes 10^{-4}$	$9.19 imes10^{-4}$	$3.490 imes 10^{-4}$	$3.75 imes 10^{-4}$
2 - 14	$9.861 imes 10^{-3}$	$1.25 imes 10^{-2}$	$4.171 imes 10^{-3}$	$4.77 imes 10^{-3}$	$1.639 imes 10^{-3}$	$1.85 imes 10^{-3}$
3 - 4	$8.916 imes10^{-1}$	$7.87 imes10^{-1}$	$8.398 imes10^{-1}$	$6.89 imes10^{-1}$	$8.177 imes10^{-1}$	$5.96 imes10^{-1}$
3 - 5	$3.383 imes 10^{-2}$	$3.61 imes 10^{-2}$	1.603×10^{-2}	1.55×10^{-2}	$7.172 imes 10^{-3}$	$6.55 imes 10^{-3}$
3 - 6	1.349×10^{-1}	1.43×10^{-1}	9.775×10^{-2}	1.19×10^{-1}	7.539×10^{-2}	1.10×10^{-1}
3 - 7	3.379×10^{0}	3.51×10^{0}	3.926×10^{0}	4.27×10^{0}	4.410×10^{0}	4.89×10^{0}
3 - 8	2.556×10^{0}	2.67×10^{0}	2.857×10^{0}	3.12×10^{0}	2.904×10^{0}	3.60×10^{0}
3 – 9	$4.186 \times 10^{\circ}$	$4.06 \times 10^{\circ}$	$4.695 \times 10^{\circ}$	$4.81 \times 10^{\circ}$	$4.780 \times 10^{\circ}$	$5.63 \times 10^{\circ}$
3 - 10	$2.088 \times 10^{\circ}$	$2.17 \times 10^{\circ}$	$2.467 \times 10^{\circ}$	$2.66 \times 10^{\circ}$	$2.677 \times 10^{\circ}$	$3.15 \times 10^{\circ}$
3 - 11	6.232×10^{-1}	6.04×10^{-1}	7.416×10^{-1}	$7.47 \times 10^{\circ}$	8.072×10^{-1}	9.00×10^{-1}
3 - 12	1.428×10^{-1}	1.90×10^{-3}	1.231×10^{-3}	1.28×10^{-1}	1.202×10^{-1}	1.09×10^{-1}
3 - 13 2 14	0.309×10^{-2}	0.50×10^{-2}	3.304×10^{-2}	4.24×10^{-2}	2.005×10^{-3}	2.75×10^{-3}
3 - 14 4 - 5	5.503×10^{-2}	5.80×10^{-2}	1.073×10^{-2} 2.671 × 10 ⁻²	1.49×10^{-2} 2.61 × 10 ⁻²	9.000×10^{-2}	1.10×10^{-2}
4 - 6	2.318×10^{-1}	2.89×10^{-1}	1.793×10^{-1}	2.01×10^{-1} 2.75 × 10 ⁻¹	1.107×10^{-1} 1.471×10^{-1}	2.74×10^{-1}
4 - 7	6.345×10^{-3}	7.20×10^{-2}	3.119×10^{-3}	7.70×10^{-3}	1.355×10^{-3}	4.21×10^{-3}
4 - 8	4.238×10^{0}	4.63×10^{0}	4.751×10^{0}	5.46×10^{0}	4.835×10^{0}	6.23×10^{0}
4 - 9	1.261×10^{1}	1.27×10^{1}	1.414×10^{1}	1.52×10^{1}	1.439×10^{1}	1.76×10^{1}
4 - 10	2.468×10^{-1}	2.79×10^{-1}	2.679×10^{-1}	$2.93 imes 10^{-1}$	2.979×10^{-1}	$3.16 imes 10^{-1}$
4 - 11	2.208×10^{0}	2.29×10^{0}	2.581×10^{0}	2.75×10^{0}	2.795×10^{0}	3.23×10^{0}
4 - 12	1.165×10^{1}	1.07×10^1	1.382×10^1	1.32×10^1	1.501×10^1	1.62×10^1
4 - 13	1.042×10^{-2}	$1.01 imes 10^{-2}$	$4.917 imes 10^{-3}$	$5.27 imes 10^{-3}$	$2.094 imes 10^{-3}$	$2.16 imes 10^{-3}$
4 - 14	$4.929 imes 10^{-2}$	$6.27 imes 10^{-2}$	$2.100 imes 10^{-2}$	$2.39 imes 10^{-2}$	$8.410 imes 10^{-3}$	$9.45 imes 10^{-3}$
5 - 6	6.396×10^{0}	6.54×10^{0}	6.502×10^0	7.69×10^{0}	6.069×10^0	8.82×10^0
5 - 7	1.214×10^{-2}	1.21×10^{-2}	6.841×10^{-3}	8.61×10^{-3}	3.973×10^{-3}	5.89×10^{-3}
5 - 8	3.337×10^{-2}	2.99×10^{-2}	1.671×10^{-2}	1.68×10^{-2}	7.776×10^{-3}	7.31×10^{-3}
5 - 9	1.169×10^{-1}	1.34×10^{-1}	8.954×10^{-2}	1.27×10^{-1}	7.004×10^{-2}	1.27×10^{-1}
5 - 10	4.295×10^{-2}	4.18×10^{-2}	1.955×10^{-2}	1.84×10^{-2}	8.514×10^{-3}	7.74×10^{-3}
5 - 11	7.151×10^{-2}	6.94×10^{-2}	3.279×10^{-2}	3.04×10^{-2}	1.459×10^{-2}	1.25×10^{-2}
5 - 12	9.742×10^{-2}	9.63×10^{-2}	4.322×10^{-2}	4.18×10^{-2}	1.770×10^{-2}	1.69×10^{-2}
5 - 13 E 14	3.939×10^{0}	4.46×10^{0}	$4.4/8 \times 10^{6}$ 1.000 × 10 ¹	5.32×10^{-1}	$4.587 \times 10^{\circ}$	0.12×10^{0}
J = 14	1.000×10^{-1}	1.04 × 10-	1.990 × 10°	∠.00 × 10 ⁻	2.101 × 10 ⁻	4.34×10^{-1}

3.2. Line Broadening Results

Two methods for line broadening calculations have been used in our work. The first is the quantum mechanical approach (Q), and the second is the semiclassical perturbation method *SCP*. To evaluate the line broadening through the second method, we need atomic parameters such as energy levels and oscillator strengths. In our SCP calculations (SCP_{SST}), we have taken atomic data of the code SST [14]. We compare our results (Q and SCP_{SST}) to the SCP calculations (SCP_{BD}) performed in [12], where atomic data have been taken from the method of Bates and Damgaard [13]. This method has been used many times with different ions, and it has been shown that the corresponding results (using the Bates and Damgaard or the SST data) are in good agreement with experimental and other theoretical results [31–33]. Many of these SCP results have been stored in the database STARK-B [34].

We have performed quantum (*Q*) and semiclassical perturbation (*SCP*) Stark broadening for 12 lines of Ar VII for electron temperature range $(2 - 50) \times 10^4$ K and at electron density $N_e = 10^{18}$ cm⁻³. We present our results in Table 4 for transitions between singlets, in Table 5 for the resonance line $3s^2$ ${}^{1}S_0-3s3p$ ${}^{1}P_1^{o}$, and in Table 6 for transitions between triplets. A comparison was made between our quantum and our semiclassical perturbation results SCP_{SST} in Tables 4 and 5. We also included the semiclassical results SCP_{BD} [12] in Table 6 in our comparison. Tables 4 and 6 show that the quantum line widths are always higher than the two semiclassical ones (SCP_{SST} and SCP_{BD}). We also found that, except for the resonance line, the ratio $\frac{Q}{SCP}$ increases and decreases with temperature. The decreasing part starts in general at $T \simeq 10^5$ K. For the resonance line $3s^2 \, {}^{1}S_0-3s3p \, {}^{1}P_1^{o}$, the ratio $\frac{Q}{SCP}$ increases with T. As per Table 5, this ratio has the same behavior as that of the other lines (increasing and after decreasing) but starts to decrease for higher temperatures ($T \simeq 10^6$ K). Table 6 shows that, in all studied cases, the SCP_{SST} widths are closer to the quantum results than the SCP_{BD} ones. The disagreement between SCP_{SST} and SCP_{BD} results is due to the difference in the source of the used atomic data.

To understand the difference between SCP and quantum calculations, we present also, in Tables 4 and 6, the contributions of elastic $(\frac{Elastic}{Total})$ and strong $(\frac{Strong}{Total})$ collisions to the SCP_{SST} line broadening. Firstly, we remark that, for $T > 10^5$ K and except the resonance line, the ratios $\frac{Elastic}{Total}$ and $\frac{Strong}{Total}$ decrease with the temperature. Secondly, we see that, for each line, as the elastic and strong collisions contributions decrease, the two results (Q and SCP) become close to each other. For electron temperature $T \le 5 \times 10^4$ K, we can detect in some cases an opposite behavior between $\frac{Elastic}{Total}$ and $\frac{Strong}{Total}$ on the one hand and the ratio $\frac{Q}{SCP}$ on the other hand. This may be due to the contributions of resonances that are dominant at low temperatures. These contributions are taken into account differently in the quantum and the semiclassical perturbative methods. Figure 1 shows the behavior of the ratios $\frac{Q}{SCP}$ and $\frac{Strong}{Total}$ with the electron temperature for the 3s3d $^{3}D_{2}$ – 3s4p $^{3}P_{1}^{o}$, 3s3p $^{3}P_{2}^{o}$ – 3s4d $^{3}D_{3}$, 3s4p $^{3}P_{2}^{o}-3s4d$ $^{3}D_{3}$, and 3s3p $^{1}P_{1}^{o}-3s4s$ $^{1}S_{0}$ transitions. In fact, the Ar VII perturbing levels i' and f' are so far from the initial (*i*) and final (*f*) levels of the considered transition ($\Delta E_{ii'}$ and $\Delta E_{ff'}$ are high) and, due to this fact, for collisions by electrons, the close collisions are important. Furthermore, with the used temperature values, the ratio $\Delta E/k_BT$ is high and consequently, the inelastic cross sections are small compared to the elastic ones that become dominant (mostly due to the close collisions). The perturbative treatment in the semiclassical approach does not correctly estimate this contribution. In that situation, it is necessary to perform more sophisticated calculations such as the quantum ones. We have shown in Elabidi et al. [11], through extensive comparisons between quantum and semiclassical Stark broadening of Ar XV lines, that the disagreement between the two results increases with the increase in strong collision contributions. Figure 2 displays the Stark widths as a function of the electron temperature at a constant electron density for two selected lines between singlets : 3s² ${}^{1}S_{0}$ -3s4p ${}^{1}P_{1}^{0}$ and 3s4p ${}^{1}P_{1}^{0}$ -3s5s ${}^{1}S_{0}$ and two lines between triplets: 3s3p ${}^{3}P_{2}^{0}$ -3s4d ${}^{3}D_{3}$, and 3s3d $^{3}D_{2} - 3s4p \, ^{3}P_{1}^{0}$.

The obtained Stark broadening parameters will be useful for the investigation and modeling of the plasma of stellar atmospheres. They will be also important for the investigation of laser-produced and inertial fusion plasmas.
Table 4. Our Stark line widths (FWHM) Q for Ar VII at electron density $N_e = 10^{18}$ cm⁻³ compared to the semiclassical results SCP_{SST} obtained using the atomic data of the code SST. $\frac{Elastic}{Total}$ and $\frac{Strong}{Total}$ are respectively the contributions of elastic and strong collisions to *SCP* line broadening. *T* is expressed in 10^4 K.

Transition	Т	Q (Å)	SCP _{SST}	Elastic Total	Strong Total	$\frac{Q}{SCP}$
	2	2.684×10^{-2}	2.380×10^{-2}	0.966	0.288	1.13
	5	$1.872 imes 10^{-2}$	$1.500 imes 10^{-2}$	0.937	0.287	1.25
3s ² ¹ S ₀ -3s3p ¹ P ₁ ⁰	10	$1.453 imes10^{-2}$	$1.070 imes 10^{-2}$	0.891	0.286	1.36
$\lambda = 585.75$ Å	20	$1.135 imes10^{-2}$	$7.640 imes10^{-3}$	0.789	0.282	1.49
	30	$9.875 imes10^{-3}$	$6.360 imes10^{-3}$	0.706	0.276	1.55
	50	$8.322 imes 10^{-3}$	$5.160 imes 10^{-3}$	0.605	0.267	1.61
	2	9.941×10^{-3}	6.070×10^{-3}	0.897	0.575	1.64
	5	6.242×10^{-3}	$2.930 imes 10^{-3}$	0.827	0.361	2.13
3s ² ¹ S ₀ -3s4p ¹ P ₁ ⁰	10	4.370×10^{-3}	2.110×10^{-3}	0.766	0.357	2.07
$\lambda = 175.5$ Å	20	$3.038 imes 10^{-3}$	$1.550 imes 10^{-3}$	0.689	0.345	1.96
	30	$2.445 imes 10^{-3}$	1.320×10^{-3}	0.641	0.334	1.85
	50	$1.848 imes 10^{-3}$	$1.090 imes 10^{-3}$	0.588	0.319	1.70
	2	6.322×10^{-2}	9.750×10^{-3}	0.944	0.155	6.48
	5	3.923×10^{-2}	$5.400 imes 10^{-3}$	0.767	0.175	7.26
3s3p ¹ P ₁ ^o -3s4s ¹ S ₀	10	2.676×10^{-2}	$3.190 imes 10^{-3}$	0.616	0.172	6.84
$\lambda = 279.2 \text{ Å}$	20	1.759×10^{-2}	2.920×10^{-3}	0.472	0.162	6.02
	30	1.344×10^{-2}	2.500×10^{-3}	0.401	0.155	5.38
	50	$9.332 imes 10^{-3}$	$2.080 imes 10^{-3}$	0.328	0.145	4.49
	2	$4.517 imes 10^{-1}$	1.360×10^{-1}	0.792	0.193	3.32
	5	$2.942 imes 10^{-1}$	$8.840 imes 10^{-2}$	0.595	0.189	3.33
3s4p ¹ P ₁ ^o -3s5s ¹ S ₀	10	$1.975 imes 10^{-1}$	6.570×10^{-2}	0.458	0.179	3.01
$\lambda = 662.6 \text{ Å}$	20	1.266×10^{-1}	5.050×10^{-2}	0.359	0.166	2.51
	30	9.671×10^{-2}	4.400×10^{-2}	0.315	0.156	2.20
	50	$6.843 imes 10^{-2}$	3.740×10^{-2}	0.274	0.144	1.83
	2	1.910×10^{-1}	$4.150 imes 10^{-2}$	0.859	0.361	4.60
	5	1.355×10^{-1}	2.650×10^{-2}	0.809	0.356	5.11
3s3d ¹ D ₂ -3s4p ¹ P ₁	10	$1.011 imes 10^{-1}$	1.930×10^{-2}	0.737	0.346	5.24
$\lambda = 489.6 \text{ Å}$	20	6.854×10^{-2}	1.430×10^{-2}	0.661	0.335	4.79
	30	5.206×10^{-2}	1.210×10^{-2}	0.611	0.323	4.30
	50	$3.602 imes 10^{-2}$	$1.010 imes 10^{-2}$	0.561	0.307	3.57
	2	1.779×10^{-2}	8.070×10^{-3}	0.825	0.085	2.20
	5	$1.119 imes 10^{-2}$	$4.750 imes 10^{-3}$	0.559	0.092	2.36
3s3p ¹ P ₁ ^o -3s5s ¹ S ₀	10	$7.845 imes 10^{-3}$	3.520×10^{-3}	0.400	0.087	2.23
$\lambda = 176.5 \text{ Å}$	20	$5.460 imes 10^{-3}$	2.710×10^{-3}	0.282	0.081	2.01
	30	$4.390 imes10^{-3}$	2.360×10^{-3}	0.229	0.076	1.86
	50	$3.303 imes 10^{-3}$	$2.000 imes 10^{-3}$	0.177	0.069	1.65

Table 5. Our Stark widths (FWHM) Q for the Ar VII $3s^2 {}^1S_0 - 3s3p {}^1P_1^o$ resonance line at electron density $N_e = 10^{18} \text{ cm}^{-3}$ compared to the semiclassical results SCP_{SST} obtained using the atomic data of the code SST. $\frac{Elastic}{Total}$ and $\frac{Strong}{Total}$ are respectively the contributions of elastic and strong collisions to SCP line broadening. T is expressed in 10^5 K.

Transition	Т	Q (Å)	SCP _{SST}	<u>Elastic</u> Total	Strong Total	$\frac{Q}{SCP}$
	0.2	$2.684 imes10^{-2}$	$2.380 imes 10^{-2}$	0.966	0.288	1.13
	0.5	1.872×10^{-2}	1.500×10^{-2}	0.937	0.287	1.25
	1	1.453×10^{-2}	1.070×10^{-2}	0.891	0.286	1.36
	2	$1.135 imes 10^{-2}$	7.640×10^{-3}	0.789	0.282	1.49
3s ² ¹ S ₀ -3s3p ¹ P ₁	3	$9.875 imes 10^{-3}$	6.360×10^{-3}	0.706	0.276	1.55
$\lambda = 585.75 \text{ Å}$	5	8.322×10^{-3}	5.160×10^{-3}	0.605	0.267	1.61
	7.5	7.271×10^{-3}	4.430×10^{-3}	0.537	0.255	1.64
	10	$6.593 imes10^{-3}$	$4.020 imes 10^{-3}$	0.494	0.245	1.64
	15	5.699×10^{-3}	$3.530 imes 10^{-3}$	0.444	0.232	1.61
	30	$4.284 imes 10^{-3}$	2.890×10^{-3}	0.389	0.211	1.48
	50	$3.327 imes 10^{-3}$	$2.530 imes 10^{-3}$	0.366	0.199	1.32

Table 6. Same as in Table 4 but we add the semiclassical results SCP_{BD} obtained in Dimitrijević et al. [12] using the atomic data from Bates and Damgaard [13]. Electron density is $N_e = 10^{18}$ cm⁻³ and *T* is expressed in 10^4 K.

Transition	Т	Q	SCP _{SST}	SCP _{BD}	$\frac{Q}{SCP_{SST}}$	$\left(\frac{Elastic}{Total}\right)$ sst	$\left(\frac{Strong}{Total}\right)_{SST}$	$\frac{Q}{SCP_{BD}}$
	2	$7.510 imes 10^{-3}$	$4.83 imes10^{-3}$	$4.21 imes 10^{-3}$	1.55	0.889	0.444	1.77
	5	$4.748 imes 10^{-3}$	$2.86 imes 10^{-3}$	$2.49 imes 10^{-3}$	1.66	0.877	0.475	1.90
3s3p ³ P ₂ ^o -3s4d ³ D ₃	10	$3.369 imes 10^{-3}$	$2.04 imes 10^{-3}$	$1.81 imes 10^{-3}$	1.65	0.861	0.472	1.85
$\lambda = 192.3 \text{ Å}$	20	2.389×10^{-3}	$1.48 imes 10^{-3}$	$1.35 imes 10^{-3}$	1.62	0.823	0.461	1.77
	30	1.967×10^{-3}	1.25×10^{-3}	$1.15 imes 10^{-3}$	1.47	0.798	0.453	1.70
	50	1.532×10^{-3}	1.02×10^{-3}	$9.51 imes 10^{-4}$	1.50	0.764	0.437	1.61
	2	1.613×10^0	$3.76 imes 10^{-1}$	$3.09 imes 10^{-1}$	4.29	0.901	0.457	5.22
	5	1.061×10^{0}	2.38×10^{-1}	1.99×10^{-1}	4.46	0.873	0.455	5.33
3s4p ³ P ₂ ^o -3s4d ³ D ₃	10	7.141×10^{-1}	1.71×10^{-1}	1.45×10^{-1}	4.18	0.824	0.448	4.92
$\lambda = 1425.9 \text{ Å}$	20	4.285×10^{-1}	1.26×10^{-1}	$1.08 imes 10^{-1}$	3.40	0.767	0.435	3.97
	30	3.073×10^{-1}	1.07×10^{-1}	9.27×10^{-2}	2.87	0.740	0.424	3.31
	50	$2.013 imes 10^{-1}$	$8.87 imes 10^{-2}$	7.77×10^{-2}	2.27	0.705	0.406	2.59
	2	7.585×10^{-2}	2.33×10^{-2}	$1.84 imes 10^{-2}$	3.26	0.928	0.406	4.12
	5	4.562×10^{-2}	1.52×10^{-2}	1.16×10^{-2}	3.00	0.878	0.397	3.93
3s3d ³ D ₂ -3s4p ³ P ₁ ⁰	10	2.896×10^{-2}	1.10×10^{-2}	8.35×10^{-3}	2.64	0.809	0.389	3.47
$\lambda = 416.0$ Å	20	1.806×10^{-2}	8.14×10^{-3}	6.12×10^{-3}	2.22	0.723	0.375	2.95
	30	1.389×10^{-2}	6.91×10^{-3}	5.18×10^{-3}	2.01	0.675	0.364	2.68
	50	1.040×10^{-2}	$5.73 imes 10^{-3}$	$4.27 imes 10^{-3}$	1.82	0.625	0.346	2.43
	2	7.605×10^{-1}	$5.36 imes10^{-1}$	$5.34 imes 10^{-1}$	1.42	0.942	0.337	1.42
	5	5.453×10^{-1}	3.44×10^{-1}	3.33×10^{-1}	1.59	0.863	0.331	1.64
3s4s ³ S ₁ -3s4p ³ P ₁ ⁰	10	4.272×10^{-1}	2.49×10^{-1}	2.40×10^{-1}	1.72	0.757	0.326	1.78
$\lambda = 1982.0 \text{ Å}$	20	3.336×10^{-1}	1.85×10^{-1}	1.79×10^{-1}	1.80	0.632	0.309	1.86
	30	2.871×10^{-1}	1.59×10^{-1}	1.53×10^{-1}	1.81	0.577	0.298	1.88
	50	2.350×10^{-1}	1.33×10^{-1}	1.28×10^{-1}	1.77	0.518	0.281	1.84
	2	1.396×10^{-2}	$8.72 imes 10^{-3}$	$5.06 imes 10^{-3}$	1.56	0.996	0.358	2.75
	5	8.893×10^{-3}	5.35×10^{-3}	2.68×10^{-3}	1.66	0.947	0.368	3.30
3s3p ³ P ₂ ^o -3s4s ³ S ₁	10	6.351×10^{-3}	3.80×10^{-3}	1.91×10^{-3}	1.67	0.860	0.368	3.31
$\lambda = 250.4$ Å	20	4.549×10^{-3}	2.78×10^{-3}	1.41×10^{-3}	1.64	0.750	0.358	3.21
	30	3.742×10^{-3}	2.35×10^{-3}	1.21×10^{-3}	1.59	0.690	0.349	3.08
	50	2.915×10^{-3}	1.94×10^{-3}	1.01×10^{-3}	1.50	0.626	0.333	2.87
	2	5.727×10^{-2}	1.69×10^{-2}	$9.11 imes 10^{-3}$	3.39	0.986	0.334	6.24
	5	2.896×10^{-2}	9.92×10^{-3}	5.85×10^{-3}	2.92	0.961	0.363	4.91
3s3p ³ P ₂ ^o -3s3d ³ D ₃	10	1.708×10^{-2}	7.02×10^{-3}	4.17×10^{-3}	2.43	0.921	0.362	4.06
$\lambda = 477.5$ Å	20	1.022×10^{-2}	5.02×10^{-3}	2.96×10^{-3}	2.04	0.830	0.357	3.43
	30	7.755×10^{-3}	4.18×10^{-3}	2.45×10^{-3}	1.86	0.764	0.353	3.14
	50	5.734×10^{-3}	3.37×10^{-3}	1.95×10^{-3}	1.70	0.684	0.343	2.92



Figure 1. Ratios $\frac{Q}{SCP_{SST}}$ (•) and $\frac{Strong}{Total}$ (•) as a function of the electron temperature for the transitions: 3s3d ${}^{3}D_{2}$ – 3s4p ${}^{3}P_{1}^{o}$ (**left up**), 3s3p ${}^{3}P_{2}^{o}$ – 3s4d ${}^{3}D_{3}$ (**right up**), 3s4p ${}^{3}P_{2}^{o}$ – 3s4d ${}^{3}D_{3}$ (**left down**), and 3s3p ${}^{1}P_{1}^{o}$ – 3s4s ${}^{1}S_{0}$ (**right down**).



Figure 2. Stark width (FWHM) W as a function of the electron temperature for transitions $3s^2 {}^{1}S_0 - 3s4p {}^{1}P_1^{o}$ (**left up**) and $3s4p {}^{1}P_1^{o} - 3s5s {}^{1}S_0$ (**right up**) at electron density $N_e = 10^{17} \text{ cm}^{-3}$, and for transitions $3s3p {}^{3}P_2^{o} - 3s4d {}^{3}D_3$ (**left down**) and $3s3d {}^{3}D_2 - 3s4p {}^{3}P_1^{o}$ (**right down**) at electron density $N_e = 10^{18} \text{ cm}^{-3}$. o: Present quantum results. •: Present SCP results. \triangle : SCP results from [12].

4. Conclusions

We have calculated in the present work quantum and semiclassical perturbation Stark broadening parameters for 12 Ar VII lines at electron temperatures from 2×10^4 to 5×10^5 K and at electron density $N_{e} = 10^{18} \text{ cm}^{-3}$. The structure and collision problem has also been treated for this ion. We have used nine configurations (1s²2s²2p⁶: 3s², 3s3p, 3p², 3s3d, 3p3d, 3s4s, 3s4p, 3s4d, and 3s5s). The structure and collisional parameters have been used in our quantum mechanical line broadening calculations. Since it is important to check their accuracy, we compared our energies to those of [22], to those obtained by the multiconfiguration Hartree-Fock method [24], and to those obtained from the AUTOSTRUCTURE code [25]. An acceptable agreement was found with the NIST results (better than 1%). We also compared our A_{ii} values with those obtained from the AUTOSTRUCTURE code [25] and with those from the SUPERSTRUCTURE code [26] using five configurations (1s²2s²2p⁶: 3s², 3s³p, 3p², 3s3d, and 3s4s). The averaged difference is about 20% with the results of [25] and about 24% with those of Christensen et al. [26]. The oscillator strengths have been compared only with the results of Christensen et al. [26], and we found an averaged agreement of about 24%. The electron-ion collision process was also studied, and collision strengths from the lowest five levels to the first 14 levels are presented at three electron energies 7.779 Ry, 13.674 Ry, and 23.336 Ry. The comparison with the collision strengths of Christensen et al. [26] indicates an agreement (averaged over the considered transitions and energies) of about 20%. The reason for the disagreement between the two results could be the difference in the number of the configurations and the difference in the partial waves taken into account in the two calculations. Stark line widths for 12 lines have been calculated using our quantum formalism. We perform also a semiclassical perturbation calculations using the structure data of the SST code. We present other semiclassical widths [12] obtained using the atomic data from Bates and Damgaard [13]. Firstly, the disagreement between the two semiclassical calculations is due to the difference in the source of atomic data. Secondly, we have shown that the disagreement between the quantum and the semiclassical widths increases with the increase in the contributions to line broadening of elastic collisions (which are mostly due to strong collisions). This is because the perturbative treatment in the semiclassical approach does not estimate very well the strong collisions. We hope that the present results can fill the lack of line broadening parameters or improve the available results for the Ar VII ion, which are of interest in the investigation and modeling of astrophysical and laboratory plasmas.

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Author Contributions: R.A. and H.E. performed the calculations; H.E., S.S.B., and M.S.D. analyzed the data; H.E. wrote the paper.

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Article Semiclassical Stark Broadening Parameters of Ar VII Spectral Lines

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Abstract: Using the semi-classical perturbation approach in the impact approximation, full width at half maximum and shift have been determined for eight spectral lines of Ar VII, for broadening by electron-, proton-, and He III-impacts. The results are provided for temperatures from 20,000 K to 500,000 K, and for an electron density of 10¹⁸ cm⁻³. The obtained results will be included in the STARK-B database, which is also in the virtual atomic and molecular data center (VAMDC).

Keywords: stark broadening; atomic data; atomic processes; line profiles; Ar VII

1. Introduction

With the development of space astronomy and satellite-born spectroscopy, trace elements—which have been without importance for astrophysics—now become increasingly important, and the corresponding data of interest for the analysis of stellar spectra. For example, spectral lines of Ar VII have been observed in the spectrum of extremely hot and massive galactic O3 If supergiant HD 93129A [1]. Additionally, Werner et al. [2] have found Ar VII lines in some of the hottest known central stars of planetary nebulae, with the effective temperatures of 95,000–110,000 K, and in (pre-) white dwarfs by analyzing high-resolution spectra from the Far Ultraviolet Spectroscopic Explorer (FUSE). We note that Stark broadening is the principal pressure broadening mechanism in such hot stars, and without the corresponding Stark broadening data, reliable analysis and modelling of high-resolution spectra are not possible.

Concerning Stark broadening parameters for Ar VII spectral lines, there is only one article where Stark broadening parameters are provided for three transitions [3]. Here, full widths at half intensity maximum (FWHM) W and shifts d for eight additional transitions have been calculated by using semiclassical perturbation method (SCP, [4,5]) for collisions of Ar VII ions with electrons, protons, and He III ions, since hydrogen and helium are the main constituents of stellar atmospheres.

2. The Impact Semiclassical Perturbation Method

The semiclassical perturbation formalism (SCP) applied here for the calculations of Stark broadening parameters, full width at half intensity maximum (FWHM-*W*), and shift of spectral line (*d*) has been formulated in [4,5], and later updates, optimisations, and innovations are presented in Sahal-Bréchot [6], Sahal-Bréchot [7], Dimitrijević et al. [8], Dimitrijević and Sahal-Bréchot [9], and Sahal-Bréchot et al. [10]. Within the frame of this method, FWHM (*W*) and shift (*d*) may be expressed by the following relation:

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$$W = N \int v f(v) dv \left(\sum_{i' \neq i} \sigma_{ii'}(v) + \sum_{f' \neq f} \sigma_{ff'}(v) + \sigma_{el} \right)$$

$$d = N \int v f(v) dv \int_{R_3}^{R_D} 2\pi \rho d\rho \sin(2\varphi_p).$$
(1)

Here, *N* is electron density, f(v) the Maxwellian velocity distribution function for electrons, ρ is the impact parameter of the incoming electron, and with i', f' are denoted the perturbing levels of the initial (*i*) and final (*f*) state. The inelastic cross-section $\sigma_{ij'}(v)$, j = i, *f* is expressed as:

$$\sum_{i'\neq i}\sigma_{ii'}(v) = \frac{1}{2}\pi R_1^2 + \int_{R_1}^{R_D} 2\pi\rho d\rho \sum_{i'\neq i} P_{ii'}(\rho, v),$$
(2)

where $P_{jj'}(\rho, v), j = i, f; j' = i', f'$ is the transition probability. The elastic cross-section is

$$\sigma_{el} = 2\pi R_2^2 + \int_{R_2}^{R_D} 2\pi\rho d\rho \sin^2 \delta + \sigma_r,$$

$$\delta = (\varphi_p^2 + \varphi_q^2)^{\frac{1}{2}}.$$
 (3)

The phase shift due to the polarization potential is φ_p (r^{-4}), and due to the quadrupolar potential φ_q (r^{-3}) (see Section 3 of Chapter 2 in Sahal-Bréchot [4]). R_1 , R_2 , R_3 , and R_D are cut-offs, defined and described in Section 1 of Chapter 3 in Sahal-Bréchot [5]. σ_r denotes the contribution of Feshbach resonances, explained in detail in [11].

A review of the theory, all approximations and details of applications is given in Sahal-Bréchot et al. [10].

3. Stark Broadening Parameter Calculations

By using Equations (1)–(3) we have calculated widths (FWHM) and shifts for eight transitions (triplets) in Ar VII spectrum. The necessary atomic energy levels have been taken from Saloman [12]. The oscillator strengths—needed for calculations—have been obtained by using the method of Bates and Damgaard [13] and the tables of Oertel and Shomo [14]. When there was no corresponding data in Oertel and Shomo [14] (for some higher levels), the needed oscillator strengths have been calculated using the method of Van Regemorter et al. [15].

In Table 1 are shown widths (FWHM) and shifts of Ar VII spectral lines broadened by electron-, proton-, and He III-impacts, for a perturber density of 10^{18} cm⁻³ and for a set of temperatures from 20,000 K to 500,000 K. The temperature range is of interest for astrophysics, laboratory plasmas, fusion research, various plasmas in technology and laser-produced plasmas. If we want to use these data for higher perturber densities, the influence of Debye screening should be checked and taken into account if needed (e.g., [16]).

The accuracy of the semiclassical perturbation method is estimated by comparison with numerous experimental data for different elements and spectral lines, and is estimated to be 20–30% (see discussion in [10]). Since Ar VII is a member of the magnesium isoelectronic sequence with relatively simple spectrum, we suppose that the error of results shown in Table 1 is not much higher than 20%.

Table 1. Electron-, proton-, and doubly charged helium-impact broadening parameters for Ar VII spectral lines, for a perturber density of 10^{18} cm⁻³ and temperatures from 20,000 to 500,000 K. The calculated wavelength of the transitions (in Å) and parameter *C* are also given. When divided by the corresponding Stark width, this parameter gives an estimate for the maximal perturber density for which the line may be treated as isolated. W_e: electron-impact full width at half maximum of intensity; *d_e*: electron-impact shift; W_{*p*:} proton-impact full width at half maximum of intensity; *d_p*: proton-impact shift; W_{*p*e++}: doubly charged helium ion-impact full width at half maximum of intensity; *d_{he++}*: doubly charged helium ion-impact shift.

Transition	T(K)	W_e (Å)	d_e (Å)	W_{H^+} (Å)	d_{H^+} (Å)	$W_{He^{++}}$ (Å)	$d_{He^{++}}$ (Å)
4s ³ S - 5p ³ P ^o	20,000	0.510E - 01	0.174E-03	0.298E-03	0.197E - 04	0.492E-03	0.272E-04
443.2 Å	50,000	0.328E-01	0.217E-03	0.898E-03	0.636E-04	0.173E-02	0.117E-03
C = 0.47E + 20	100,000	0.243E-01	0.158E-03	0.148E - 02	0.129E-03	0.291E-02	0.249E-03
	200,000	0.186E - 01	0.215E-03	0.203E-02	0.223E-03	0.403E - 02	0.444E - 03
	300,000	0.162E - 01	0.211E-03	0.220E-02	0.275E-03	0.439E - 02	0.550E-03
	500,000	0.139E-01	0.227E-03	0.242E-02	0.352E-03	0.482E - 02	0.709E-03
3p ³ P ^o - 4s ³ S	20,000	0.490E - 02	-0.779E-03	0.744E - 06	0.135E - 04	0.129E-05	0.186E - 04
250.4 A	50,000	0.270E - 02	0.615E - 04	0.642E - 05	0.432E - 04	0.124E - 04	0.792E - 04
C = 0.32E + 20	100,000	0.191E - 02	0.113E-03	0.295E - 04	0.835E - 04	0.579E - 04	0.161E - 03
	200,000	0.142E - 02	0.131E-03	0.819E - 04	0.133E-03	0.163E-03	0.265E - 03
	300,000	0.121E - 02	0.143E-03	0.113E-03	0.162E - 03	0.227E-03	0.323E-03
	500,000	0.101E-02	0.136E-03	0.173E-03	0.196E-03	0.347E-03	0.396E-03
$3p^{3}P^{0} - 3d^{3}D$	20,000	0.911E - 02	-0.564E-03	0.361E - 05	-0.107E-05	0.598E - 05	-0.148E - 05
477.5 Å	50,000	0.585E - 02	-0.436E - 04	0.148E - 04	-0.348E - 05	0.282E - 04	-0.639E - 05
C = 0.48E + 21	100,000	0.417E - 02	-0.288E - 04	0.396E - 04	-0.740E - 05	0.767E - 04	-0.143E-04
	200,000	0.297E - 02	-0.165E-04	0.900E - 04	-0.150E - 04	0.176E - 03	-0.298E - 04
	300,000	0.245E - 02	-0.134E-04	0.129E - 03	-0.220E - 04	0.253E - 03	-0.438E - 04
	500,000	0.195E-02	-0.237E-04	0.177E-03	-0.333E-04	0.349E-03	-0.669E - 04
3p ³ P ^o - 4d ³ D	20,000	0.421E - 02	-0.445E-03	0.671E - 05	0.621E - 05	0.111E - 04	0.858E - 05
192.3 Å	50,000	0.249E - 02	0.896E - 05	0.263E - 04	0.200E - 04	0.506E - 04	0.366E - 04
C = 0.94E + 19	100,000	0.181E - 02	0.259E - 04	0.589E - 04	0.394E - 04	0.115E - 03	0.760E - 04
	200,000	0.135E - 02	0.428E - 04	0.100E - 03	0.646E - 04	0.199E - 03	0.128E - 03
	300,000	0.115E - 02	0.345E - 04	0.127E - 03	0.782E - 04	0.252E - 03	0.156E - 03
	500,000	0.951E-03	0.379E-04	0.155E-03	0.971E-04	0.308E-03	0.196E-03
$4p^{3}P^{0} - 4d^{3}D$	20,000	0.309	-0.849E-02	0.811E-03	0.262E - 03	0.134E - 02	0.362E - 03
1425.9 Å	50,000	0.199	0.324E - 03	0.292E - 02	0.845E - 03	0.562E - 02	0.155E - 02
C = 0.52E + 21	100,000	0.145	0.336E-03	0.577E - 02	0.169E - 02	0.113E-01	0.327E - 02
	200,000	0.108	0.126E - 02	0.877E - 02	0.286E - 02	0.173E - 01	0.569E - 02
	300,000	0.927E - 01	0.595E - 03	0.106E - 01	0.346E - 02	0.211E - 01	0.691E - 02
	500,000	0.777E-01	0.645E-03	0.120E-01	0.440E-02	0.237E-01	0.886E-02
$3d^{3}D - 4p^{3}P^{0}$	20,000	0.184E - 01	-0.111E-02	0.352E - 04	0.757E - 05	0.584E - 04	0.105E - 04
416.0 A	50,000	0.116E - 01	0.474E - 04	0.133E-03	0.245E - 04	0.256E - 03	0.450E - 04
C = 0.87E + 20	100,000	0.835E - 02	0.115E-03	0.286E-03	0.513E-04	0.560E-03	0.991E - 04
	200,000	0.612E-02	0.105E-03	0.469E-03	0.946E-04	0.926E-03	0.188E-03
	300,000	0.518E-02	0.121E-03	0.577E-03	0.126E-03	0.114E-02	0.251E-03
	500,000	0.427E-02	0.140E-03	0.679E-03	0.161E-03	0.135E-02	0.323E-03
$3d^{3}D - 5p^{3}P^{0}$	20,000	0.139E-01	-0.555E-03	0.900E-04	0.185E-04	0.148E-03	0.256E-04
240.6 A	50,000	0.868E-02	0.137E-03	0.271E-03	0.587E-04	0.523E-03	0.108E-03
C = 0.14E + 20	100,000	0.642E-02	0.162E-03	0.445E-03	0.109E-03	0.877E-03	0.211E-03
	200,000	0.490E-02	0.190E-03	0.616E-03	0.166E-03	0.122E-02	0.331E-03
	300,000	0.425E-02	0.199E-03	0.669E-03	0.204E-03	0.133E-02	0.406E-03
	500,000	0.362E-02	0.199E-03	0.739E-03	0.237E-03	0.147E-02	0.479E-03
$4d^{3}D - 5p^{3}P^{0}$	20000	0.243	-0.224E-03	0.163E-02	0.134E-03	0.269E-02	0.185E-03
952.0 A	50,000	0.158	0.166E-02	0.477E-02	0.431E-03	0.923E-02	0.790E-03
C = 0.22E + 21	100,000	0.118	0.174E-02	0.767E-02	0.856E-03	0.151E-01	0.165E-02
	200,000	0.902E-01	0.184E-02	0.104E-01	0.142E-02	0.207E-01	0.283E-02
	300,000	0.786E-01	0.221E-02	0.113E-01	0.172E-02	0.224E-01	0.343E-02
	500,000	0.6/2E-01	0.208E-02	0.124E - 01	0.21/E - 02	0.246E-01	0.43/E-02

Since the wavelengths in Table 1 are calculated ones, they are different from experimental wavelengths. However, we notice that they are correct in angular frequency units, because in such

a case, relative and not absolute positions of energy levels are important for calculations. For the transformation of the Stark widths from Å-units to angular frequency units, the following formula can be used:

$$W(\mathring{A}) = \frac{\lambda^2}{2\pi c} W(s^{-1}) \tag{4}$$

where *c* is the speed of light. For the correction of widths and/or shifts for the difference between calculated and experimental wavelength, one can use the expression:

$$W_{cor} = \left(\frac{\lambda_{exp}}{\lambda}\right)^2 W.$$
(5)

The corresponding expressions for the shifts are analogous to Equations (4) and (5). Here, W_{cor} denotes the corrected width, while λ_{exp} is the experimental wavelength, λ the calculated wavelength, and W the width from Table 1.

Dividing the parameter C [17] from Table 1 by the corresponding full width at half maximum, one obtains the maximal perturber density for which the line may be treated as isolated.

The obtained Ar VII Stark broadening parameters shown in Table 1 will be implemented in the STARK-B database [18,19]. This database contains Stark widths and shifts needed first of all for the investigations, modelling, and diagnostics of the plasma of stellar atmospheres, but also for diagnostics of laboratory plasmas and investigation of laser produced, inertial fusion plasma, and for plasma technologies.

The STARK-B database is one of the databases included in the virtual atomic and molecular data center—VAMDC [20,21], created in order to enable more effective search and mining of atomic and molecular data which are in different databases. Databases with atomic and molecular data which are in VAMDC—including STARK-B—can be accessed and searched through the VAMDC portal: http://portal.vamdc.org/.

4. Conclusions

The semiclassical perturbation calculation of Stark broadening parameters, widths, and shifts for spectral lines broadened by collisions of Ar VII ions with electrons, protons, and doubly charged helium ions have been performed for eight multiplets of Ar VII. The obtained values of Stark broadening parameters will be implemented in the STARK-B database—one of the databases included in the virtual atomic and molecular data center (VAMDC). Since Stark broadening data for Ar VII spectral lines considered here do not exist in the literature, we hope that they will be of interest for the relevant problems in astrophysical, laboratory, laser-produced, inertial fusion, and technological plasmas.

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Article Stark Widths of Ar II Spectral Lines in the Atmospheres of Subdwarf B Stars

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Abstract: Stark broadening parameters are of interest for many problems in astrophysics and laboratory plasmas investigation. Ar II spectral lines are observed in many kinds of stellar atmospheres such as the atmospheres of B-Type stars and subdwarf B stars. In this work, we present theoretical Stark widths for Ar II spectral lines. We use the impact semiclassical perturbation approach. Our results are compared with the available experimental values. Finally, the importance of the Stark broadening mechanism is studied in atmospheric conditions of subdwarf B stars.

Keywords: stark broadening; atomic data; subdwarfs B stars; Ar II

1. Introduction

Argon in different ionization stages is important for modelling and investigating stellar atmospheres. For example, Werner et al. [1] used Ar VII λ = 1063.55 Å spectral line for abundance determination of argon in the extremely hot helium-rich white dwarf PG 1034 + 001. In [2], the discovery of argon in hot evolved stars and white dwarfs has been reported. Rauch et al. [3] have identified Ar VI absorption lines in the spectrum of a hydrogen-rich central star using high-resolution, high signal to noise observations obtained with Far Ultraviolet Spectroscopic Explorer (FUSE) and Space Telescope Imaging Spectrograph (STIS) aboard the Hubble Space Telescope (HST). In [4], the Ar III λ = 1002.097 Å spectral lines were also observed by O'Toole and Heber [5] in the spectra of subdwarf B stars. Ar III spectral lines were also observed by O'Toole and Heber [5] in the spectra of subdwarfs. Keenan et al. [7] have determined the abundance of argon from stellar absorption lines of Ar II in the optical spectra of main-sequence early-type stars. Recently, Ar II spectral lines were observed by kupfer et al. [8] in extreme helium stars.

Stark broadening is an important broadening mechanism for modelling and investigating stellar atmospheres and for the determination of abundance of chemical elements. For example, the importance of the Stark broadening effect in stellar atmospheres of A and B type stars was studied by Popović et al. [9] and Simić et al. [10]. Based on Si VI lines, Hamdi et al. [11] demonstrated that the Stark broadening mechanism is dominant in broad regions in the studied atmospheres of DO white dwarfs. The influence of Stark broadening in DO white dwarf atmospheres was also

studied by Dimitrijević et al. [12] using Xe VIII spectral lines and similar conclusions were found. Hamdi et al. [13] studied the importance of Stark broadening in the atmospheres of subdwarf B stars using Ar III spectral lines. It was demonstrated that this mechanism is important, especially for atmospheres with high values of log g.

In this paper, we have reported Stark widths of 34 spectral lines of Ar II ions belonging to the 3d–4p transition array. Calculations were performed using the semiclassical perturbation approach (SCP) in impact approximation [14,15]. Energy levels and oscillator strengths needed as input parameters for Stark width calculation were calculated using the Hartree–Fock relativistic approach using Cowan code [16]. We use an atomic model including 24 configurations. Our results are compared with the experimental values. In the last section of this paper, we have studied the importance of the Stark broadening mechanism in the atmospheres of subdwarf B Stars. Stark and Doppler widths are compared as a function of the temperature of the atmospheric layers and as a function of optical depth.

2. The Impact Semiclassical Perturbation Method

In the semiclassical perturbation approach, the full width at half maximum (W) can be expressed in terms of the inelastic cross-section and elastic processes as:

$$W = N \int v f(v) dv \left(\sum_{i' \neq i} \sigma_{ii'}(v) + \sum_{f' \neq f} \sigma_{ff'}(v) + \sigma_{el} \right)$$

where *N* is the density of colliding perturbers, f(v) is the Maxwell distribution of the relative atom-perturber velocity v, $\sigma_{ii'}$ and $\sigma_{ff'}$ are the inelastic cross-sections between the initial level *i* (resp. *f* the final level) and the perturbing levels *i'* (resp. *f'*) of *i* – *f* transition. $\sigma_{el}(v)$ represents the contribution of elastic collisions and includes Feshbach resonances when ion–electron impacts are studied.

The main input data used in the semiclassical perturbation calculation of Stark broadening parameters are energy levels and oscillator strengths. The set of atomic data needed for the semiclassical method is relatively large. In this paper, the set of energy levels and oscillator strengths needed for the SCP calculation of Stark widths is calculated using the multiconfiguration Hartree–Fock method with relativistic correction (HFR) [16]. We use an atomic model including 24 configurations: $3s^2 3p^5$; $3s^2 3p^4$ nl (nl = 4p, 4f, 5p, 5f, 6p, 6f, 6h, 7p, 7f, 7h) (odd parity) and $3s 3p^6$; $3s^2 3p^4 n'l'$ (n'l' = 3d, 4s, 4d, 5s, 5d, 5g, 6s, 6d, 6g, 7s, 7d, 7g) (even parity).

In order to introduce a correction to the widths due to the difference between calculated and experimental wavelengths, we have used Equation (8) of Hamdi et al. [17].

3. Stark Widths

Our Stark widths calculated as described above are presented in Table 1 along with the experimental results of [18–23]. Table 1 is organized in the following way: In the first five columns of Table 1, we give the transition array, transition, wavelength, electron temperature (T) and electron density (N_e). In column 6 of Table 1, we show the experimental Stark widths (W_m). In columns 7 and 8, we present our electron impact Stark widths (W_i) and ion impact Stark widths (W_e) calculated using the semiclassical perturbation approach as described in Section 2. Ionic Perturbers are Ar^+ ions. The ratios (W_m/W) between the experimental and our calculated Stark widths are shown in column 9, where $W = W_e + W_i$ is the total width. Finally, in the last two columns, we give the accuracies and the references of the experimental results. For the estimates, we use code letters used by Konjević et al. [24] which indicate the following:

- A = uncertainties within 15%
- B+ = uncertainties within 23%
- B = uncertainties within 30%

- C+ = uncertainties within 40%
- C = uncertainties within 50%
- D = uncertainties larger than 50%

Generally, for the (³P) 3d–(³P) 4p transitions, our Stark widths agree well with the experimental results. The difference between our Stark widths and Pellerin et al. [20] ones does not exceed 31% except for two transitions: ${}^{4}D_{5/2}-{}^{2}D_{5/2}^{0}$ ($\lambda = 3808.57$ Å) and ${}^{4}P_{5/2}-{}^{4}S_{3/2}^{0}$ ($\lambda = 3499.48$ Å) for which the differences are 39% and 56% respectively. We note that for the transition ${}^{4}D_{5/2}-{}^{2}D_{5/2}^{0}$ ($\lambda = 3808.57$ Å), the ratio $\frac{W_{m}}{W}$ is equal to 0.87 when we compare with Djurović et al. [23]. Our results underestimate the experimental values of Pellerin et al. [20] for four transitions. For the other sixteen transitions, the values of Pellerin et al. [20] are overestimated.

On average, our Stark width agrees with Aparicio et al. [21] and Djurović et al. [23] within 20%. The largest difference found with Djurović et al. [23] is for ${}^{4}D_{5/2} - {}^{4}S_{3/2}^{o}(\lambda = 3499.48 \text{ Å})$ transition for which the ratio $\frac{W_{m}}{W}$ is equal 1.42. We note here that the value is measured with an error between 30% and 50%. The values of Aparicio et al. [21] are underestimated for seven transitions and overestimated for the other eighteen transitions.

Our Stark widths are compared with Iglesias et al. [22] for two transitions. Results of Iglesias et al. [22] are given for three values of electron densities: 4.5×10^{17} cm⁻³, 7.2×10^{17} cm⁻³ and 9.0×10^{17} cm⁻³. Comparing with Iglesias et al. [22], the average difference is 21% but a large difference is found for the transition ${}^{4}D_{5/2} - {}^{4}D_{5/2}^{o}$ ($\lambda = 3968.6$ Å) for the value measured at 7.20×10^{17} cm⁻³ electron density. In fact, for this value, the ratio $\frac{W_{m}}{W}$ is equal to 0.62. On the other hand, for the values measured at 4.5×10^{17} cm⁻³ and 9.0×10^{17} cm⁻³ electron densities, the ratio $\frac{W_{m}}{W}$ is equal 0.80.

Our Stark widths are compared with Dzierzega and Musiol [19] for the transition ${}^{4}D_{7/2} - {}^{4}D_{7/2}^{0}$ ($\lambda = 4013.86$ Å). Dzierzega and Musiol [19] have measured Stark widths for different values of temperature and electron densities. All our Stark widths overestimate those of Dzierzega and Musiol [19]. The average ratio $\frac{W_{m}}{W}$ is equal to 0.74. Better agreement with Dzierzega and Musiol [19] ($\frac{W_{m}}{W} = 0.96$) is found for the width measured at the temperature 12,200 K and the electron density 0.74 × 10¹⁷ cm⁻³. The largest disagreement is found for the width measured at the temperature 11,520 K and the electron density 1.79×10^{17} cm⁻³.

In Figure 1, we present our electron impact Stark width as a function of temperature for the interval from 5000 to 60,000 K along with the experimental values of [18–23] for the transition (³P) 3d ⁴D_{7/2}–(³P) 4p ⁴D_{7/2}⁰ (λ = 4013.86 Å). All the experimental results are normalized to an electron density of 10¹⁷ cm⁻³. Figure 1 shows that our Stark widths are close to the experimental results. The closest values to our widths are those of Djurović et al. [23] and of Dzierzega and Musiol [19] measured at the temperature of 12,200 K and the electron density of 0.74 × 10¹⁷ cm⁻³. Figure 1 shows also that our Stark widths overestimate the result of Iglesias et al. [22] measured at the electron density equal to 9.0 × 10¹⁷ cm⁻³ and underestimate values of Iglesias et al. [22] measured at the electron densities equal to 4.5 × 10¹⁷ cm⁻³ and 7.2 × 10¹⁷ cm⁻³.

Table 1. Our electron impact Stark widths (FWHM) (W_e) and ion impact Stark width (W_i) calculated using the semiclassical perturbation (SCP) approach in impact approximation [14,15] compared with experimental Stark widths (W_m). Transitions, wavelengths, electron temperature (T) and electron density (N_e) are also given. All wavelengths are taken from the NIST database [25].

Transition Array	Transition	λ (Å)	$T \; (10^3 \; K)$	$N_e (10^{17} \text{ cm}^{-3})$	W_m	W_e (pm)	W _i (pm)	W_m/W	Acc.	Ref.
(³ P) 3d- (³ P) 4p	${}^{4}\text{D}_{7/2} - {}^{4}\text{P}_{5/2}^{0}$	4400.99	22.0	1.00	34.3	25.2	3.78	1.18	В	[20]
	${}^{4}D_{5/2} - {}^{4}P_{3/2}^{0}$	4371.33	22.0	1.00	32.4	25.0	3.77	1.13	В	[20]
	${}^{4}D_{5/2} - {}^{4}P_{5/2}^{0}$	4431.00	22.0	1.00	28.6	25.3	3.84	0.98	В	[20]
			18.4-26.5	1.00	31.3	26.8-24.0	3.71-3.93	1.02 - 1.12	В	[21]
	${}^{4}D_{3/2} - {}^{4}P^{o}_{3/2}$	4400.10	22.0	1.00	30.7	25.1	3.82	1.06	В	[20]
	${}^{4}D_{1/2} - {}^{4}P_{1/2}^{0}$	4352.20	22.0	1.00	35.0	24.9	3.78	1.22	В	[20]
	${}^{4}D_{3/2} - {}^{4}P_{5/2}^{0}$	4460.56	22.0	1.00	26.6	25.4	3.89	0.91	В	[20]
	4D 4D0	4420.01	18.4-26.5	1.00	36.8	27.0-24.1	3.78-4.00	1.19-1.31	В	[21]
	$D_{1/2} - T_{3/2}$	4420.91	18 4 24 5	1.00	24.4	23.0	276 207	1.00 1.20	Б Ст	[20]
	${}^{4}D_{7} = {}^{4}D_{7}^{0}$	4013.86	26.0	1.00	34.0	38.3	6.04	0.77	B	[18]
	27/2 27/2	1010100	10.88	2.03	46.7	60.4	5.89	0.70	B+	[19]
			11.52	1.79	35.3	52.1	5.29	0.62	B+	[19]
			12.20	0.74	22.4	21.0	2.04	0.96	B+	[19]
			13.03	1.10	25.6	30.5	3.37	0.76	B+	[19]
			13.88	1.39	26.7	37.6	4.29	0.64	B+ B	[19]
			18.4-26.5	1.00	21.5	24.4-21.6	3.26-3.45	0.77-0.86	B+	[20]
			43.0	4.50	100	85.7	16.2	0.98	В	[22]
			43.0	7.20	160	137	25.9	0.98	В	[22]
			43.0	9.00	160	171	32.3	0.79	В	[22]
	40 400	20(0.2)	22.0	1.00	23.34	23.0	3.36	0.86	В	[23]
	$D_{5/2} - D_{5/2}^{o}$	3968.36	22.0 19.4.06 E	1.00	23.6	22.6	3.33	0.91	B P.	[20]
			43.0	4 50	29.4	24.0-21.4 84.8	3.24-3.43	0.79	B+	[21]
			43.0	7.20	100	136	25.7	0.62	B	[22]
			43.0	9.00	160	169	32.1	0.80	В	[22]
	⁴ D _{3/2} - ⁴ D ⁰ _{3/2}	3914.77	22.0	1.00	20.7	21.7	3.30	0.83	В	[20]
			18.4-26.5	1.00	20.9	23.1-20.6	3.20-3.40	0.79 - 0.87	B+	[21]
	45 450		22.0	1.00	18.39	21.7	3.30	0.74	В	[23]
	$^{4}D_{7/2} - ^{4}D_{5/2}^{0}$	3944.27	22.0	1.00	22.0	22.5	3.29	0.85	B	[20]
			18.4-26.5 22.0	1.00	24.0	23.8-21.2	3.19-3.38	0.90-0.98	D+ A	[21]
	${}^{4}\text{D}_{2}/_{2} - {}^{4}\text{D}_{2}^{0}/_{2}$	3875.26	22.0	1.00	19.2	22.0	3.26	0.76	B	[20]
	-3/2 -1/2		18.4-26.5	1.00	27.5	23.4-20.6	3.17-3.36	1.04-1.15	B+	[21]
			22.0	1.00	17.62	22.0	3.26	0.70	В	[23]
	${}^{4}D_{5/2} - {}^{4}D_{7/2}^{0}$	4038.80	22.0	1.00	24.1	23.2	3.41	0.91	В	[20]
			18.4-26.5	1.00	29.9	24.6-21.9	3.31-3.51	1.07 - 1.18	В	[21]
	40 400	2002.05	22.0	1.00	24.05	23.2	3.41	0.90	C	[23]
	$D_{3/2} - D_{5/2}^{o}$	3992.05	22.0 19.4.06 E	1.00	21.9	22.9	3.38	0.83	B P.	[20]
	$^{4}D_{e} = ^{4}D^{0}$	3931 24	18.4-26.5 22.0	1.00	32.4 19.1	24.3-21.7	3.2/-3.4/	0.76	B+	[21]
	${}^{4}D_{7/2} - {}^{2}D_{3/2}^{0}$	3786.38	22.0	1.00	20.3	22.2	3.17	0.79	B	[20]
	27/2 25/2	0700.00	18.4-26.5	1.00	23.7	23.8-21.3	3.08-3.27	0.88-0.96	B+	[21]
			22.0	1.00	23.49	22.4	3.17	0.92	А	[23]
	⁴ D _{5/2} - ² D ⁰ _{5/2}	3808.57	22.0	1.00	18.7	22.8	3.19	0.72	В	[20]
			22.0	1.00	22.53	22.8	3.19	0.87	С	[23]
	⁴ D _{5/2} - ⁴ S ^o _{3/2}	3499.48	22.0	1.00	29.03	17.5	2.90	1.42	С	[23]
	${}^{4}F_{7/2} - {}^{2}D_{5/2}^{0}$	6243.12	18.4-26.5	1.00	60.3	68.7-61.4	8.64-9.14	0.78-0.85	B+	[21]
	${}^{4}F_{5/2} - {}^{2}D_{3/2}^{0}$	6138.66	18.4-26.5	1.00	58.6	67.0-59.6	8.57-9.07	0.78-0.85	B+	[21]
	$F_{5/2} - D_{5/2}^{0}$	6399.21	18.4-26.5	1.00	49.5	72.4-64.6	9.07-9.59	0.61-0.68	B+	[21]
	$^{-P_{1/2}-D_{3/2}^{\circ}}$	6808.55	22.0 19.4.06 E	1.00	/5.3	77.1	10.9	0.86	D D	[20]
	$r_{3/2} - r_{3/2}$	6666 26	18.4-20.5	1.00	62.6	74.9-07.1	10.7.11.2	0.75-0.80	D+ B	[21]
	$^{1}_{2P_{1/2}}$	6437.60	10.4-20.3	1.00	65.0	70.3-02.8 65.6-58.6	9.98-10.5	0.77-0.04	B+	[21] [21]
	$^{2}P_{2} = ^{2}S^{0}$	6483.08	18.4-26.5	1.00	65.2	71 1-64 2	10 35-10 9	0.80-0.87	B+	[21]
	${}^{2}P_{1/2} - {}^{2}S_{0}^{0}$	6103.54	22.0	1.00	65.3	59.2	9,36	0.95	В	[20]
	- 1/2 01/2	5100.01	18.4-26.5	1.00	61.3	62.6-56.3	9.14-9.65	0.85-0.93	B+	[21]
	${}^{4}P_{3/2} - {}^{4}S^{0}_{3/2}$	7380.43	18.4-26.5	1.00	76.7	87.6-76.1	13.2-13.9	0.76-0.85	B+	[21]
	${}^{4}P_{1/2} - {}^{4}S_{3/2}^{0}$	7233.54	22.0	1.00	61.9	83.2	13.0	0.64	C+	[20]
	4F3/2-4D3/2	6756.55	18.4-26.5	1.00	62.8	73.5-65.5	9.89-10.5	0.75-0.83	B+	[21]
	${}^{4}F_{5/2} - {}^{4}D_{5/2}^{0}$	6863.54	18.4-26.5	1.00	55.4	76.5-68.1	9.98-10.56	0.64-0.70	А	[21]
	4F7/2-4D5/2	6684.29	18.4-26.5	1.00	61.6	72.5-64.6	9.50-10.1	0.75-0.82	B+	[21]
	4F7/2-4D7/2	6886.61	18.4-26.5	1.00	69.5	76.2-67.2	9.88-10.5	0.81-0.89	А	[21]
	${}^{4}F_{9/2} - {}^{4}D_{7/2}^{0}$	6643.70	18.4-26.5	1.00	71.8	72.3-64.0	9.20-9.70	0.88-0.97	B+	[21]



Figure 1. Electron impact Stark width (FWHM) obtained using the semiclassical perturbation approach [14,15] for (³P) 3d ${}^{4}D_{7/2}$ -(³P) 4p ${}^{4}D_{7/2}^{0}$ (λ = 4013.86 Å) spectral lines as a function of electron temperature compared with experimental results. Electron density is 10¹⁷ cm⁻³.

4. Stark Broadening Effect in sdB Stars

Subdwarf B stars are low-mass (roughly half a solar mass) helium burning stars with extremely thin hydrogen envelopes. They behave as helium main sequence stars. The sdB stars have a high effective temperature (20,000 K \leq T_{eff} \leq 40,000 K) and gravities (log $g \simeq$ 5–6) (see e.g., [26]). Ar II spectral lines are observed in the atmospheres of sdB stars. For example, Heber and Edelmann [6] have reported abundance of argon in B subdwarfs using Ar II spectral lines.

Beside broadening by collisions with electrons (Stark broadening), Doppler broadening is also important in stellar atmospheres. The intensity distribution function is Lorentzian in the case of Stark broadened lines and Gaussian in the case of Doppler broadened lines. In order to study the importance of Stark broadening in the atmospheric conditions of subdwarf B stars, we have compared Stark and Doppler widths for Ar II (³P) 3d ⁴D_{5/2}–(³P) 4p ⁴P^o_{3/2} (λ = 4371.33 Å) spectral lines. We used the atmospheric models of Jeffery et al. [27] which are plane-parallel line-blanketed model atmospheres for hot stars in local thermal, radiative and hydrostatic equilibrium. The considered atmospheres have the following composition: 0.001 helium, 0.99741 hydrogen and 0.00047 carbon and nitrogen.

In Figures 2 and 3, we present Stark and Doppler widths for Ar II (³P) 3d ⁴D_{5/2}–(³P) 4p ⁴P_{0/2}^o ($\lambda = 4371.33$ Å) spectral lines as a function of the temperature of the atmospheric layers and as a function of optical depth respectively. Figure 2 shows that for the atmospheres with log *g* = 6.00 and log *g* = 5.75, Stark width become larger than Doppler width from the atmospheric layer with temperature T = 55,000 K and T = 60,000 K respectively. For the atmosphere with log *g* = 5.50, Stark width is comparable to Doppler width only for the dipper layers of the atmosphere. For the atmospheres with log *g* = 5.25 and log *g* = 5.00, Stark width is dominated by Doppler width in all atmospheric layers. Due to the different behavior of Gaussian and Lorentzian distributions, Stark broadening may be important in the line wings even when Doppler width is larger than Stark width.



Figure 2. Stark and Doppler widths for Ar II (³P) 3d ${}^{4}D_{5/2}$ –(³P) 4p ${}^{4}P_{3/2}^{o}$ (λ = 4371.33 Å) spectral lines as a function of atmospheric layer temperature. Stark widths are shown for five values of model gravity log g = 5 - 6, $T_{eff} = 22,000$ K.



Figure 3. Stark and Doppler widths for Ar II (³P) 3d ${}^{4}D_{5/2}$ –(³P) 4p ${}^{4}P_{3/2}^{o}$ (λ = 4371.33 Å) spectral lines as a function of optical depth. Stark widths are shown for five values of model gravity log g = 5 - 6, $T_{eff} = 22,000$ K.

5. Conclusions

Using the Hartree–Fock approach with relativistic corrections for the calculations of energy levels and oscillator strengths, and the semiclassical perturbation approach in impact approximation, we have determined Stark widths for 34 spectral lines of Ar II ion. All studied lines belong to the 3d–4p transition array. The comparison of our results shows that they are, generally, in good agreement with the experimental values. Our study will be extended to other transition arrays. Stark shift will also be studied. Our investigation on the importance of Stark broadening in the atmospheres of subdwarf B stars, shows that Stark broadening is an important broadening mechanism for the atmospheres with high surface gravity. So, Stark widths obtained here may be useful for modeling and investigating those kind of stars.

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Author Contributions: Rafik Hamdi carried out the calculations. All the authors were involved in the discussions of results and in writing the manuscript.

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Article Stark Broadening of Se IV, Sn IV, Sb IV and Te IV Spectral Lines

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Abstract: Stark broadening parameters, line width and shift, are needed for investigations, analysis and modelling of astrophysical, laboratory, laser produced and technological plasmas. Especially in astrophysics, due to constantly increasing resolution of satellite borne spectrographs, and large terrestrial telescopes, data on trace elements, which were previously insignificant, now have increasing importance. Using the modified semiempirical method of Dimitrijević and Konjević, here, Stark widths have been calculated for 2 Se IV, 6 Sn IV, 2 Sb IV and 1 Te IV transitions. Results have been compared with existing theoretical data for Sn IV. Obtained results will be implemented in the STARK-B database, which is also a part of Virtual atomic and molecular data center (VAMDC).

Keywords: Stark broadening; line profiles; atomic data

1. Introduction

With the development of satellite-born spectrographs, the significance of trace elements, which have been without any particular importance for investigation of stellar spectra before the development of space astronomy, is becoming more and more increasing. For example, Rauch et al. [1] underlined that accurate Stark broadening data for spectral lines of as much as possible large number of atoms and ions "are of crucial importance for sophisticated analysis of stellar spectra by means of NLTE model atmospheres". High resolution spectra obtained from space born instruments contain different lines of trace elements and the corresponding Stark broadening data are important for their analysis and synthesis. But such data are also very useful for laboratory plasma diagnostics and for investigation and modelling of various plasmas in technology, inertial fusion, as well as for research of laser produced plasmas.

For example, Selenium (Se), which was previously without astrophysical significance, has been detected in the atmospheres of Am star ρ Pup [2], in cool DO white dwarfs [3,4], and Se III emission has been identified in the planetary nebula (PN) NGC5315 [5]. Werner et al. [6] found Se, Sn and Te in the spectra of RE0503-289, a helium rich DO white dwarf. Presence of selenium in the stellar spectra is a confirmation that in subphotosperic layers, where Stark broadening is dominant broadening mechanism [7] it exists in various ionization stages. Consequently, the corresponding Stark broadening parameters for Se in various ionization stages, are useful for theoretical consideration and modelling of such layers, as well as for the calculation of radiative transfer in such conditions. Moreover, in DO white dwarfs, where effective temperatures are within the interval 45,000 K–120,000K, Stark broadening is usually larger than or comparable to Doppler broadening [8,9]. At such temperatures selenium is

mainly ionized and if one ionization stage is observed, certainly others also exist and will be observed in the future with new generations of space telescopes and large telescopes on the Earth. In spite of the fact that Stark broadening data for various ionization stages are needed for interpretation, analysis and synthesis of selenium spectral lines in astrophysical spectra, exist only data for Se I [10,11].

Tin (Sn) is also present in stellar spectra. As an example, Sn lines have been observed in the spectra of A-type stars [12] and in cool DO white dwarfs [3,4]. They mark the hot end of the so-called DB gap, which corresponds to an interval in effective temperatures from 45,000 K to 30,000 K [4]. For a wider analysis of the tin presence in stellar spectra see references in Alonso-Medina and Colón [13]. It is worth to note that Proffitt et al. [14] used the 1313.5-Å resonance line of Sn IV in order to determine the tin abundance of the early B main-sequence star, AV 304, in the Small Magellanic Cloud. Alonso-Medina and Colón [13] report also that Sn plasmas are a candidate for the extreme ultraviolet light source for next-generation microlithography [15,16]. Stark broadening of Sn II and Sn III lines has been measured by Kieft et al. [17], in order to investigate the pinched discharge plasmas in tin vapor as candidates for application in future semiconductor lithography. Stark broadening of seven Sn III spectral lines has been also measured by Djeniže [18]. Concerning the spectral lines of Sn IV, experimentally determined Stark broadening parameters of nine spectral lines have been published by Djeniže [18], and Burger et al. [19] measured again five of them. Results of theoretical calculations of Stark broadening parameters of Sn III lines can be found in [13,17,20]. For Sn IV, there is only one paper [21] with Stark widths and shifts for 66 spectral lines, calculated using semiempirical method of Griem [22] with atomic matrix elements obtained with the relativistic Hartree-Fock method and configuration interaction in an intermediate coupling scheme, by using the Cowan [23] approach and the COWAN code.

Spectral lines of antimony (Sb) are also present in stellar spectra. For example, very strong absorption Sb II spectral lines are observed in the spectrum of HgMn star HR7775 where Stark broadening can not be neglected [24]. Besides the astrophysical importance, antimony is also significant in thin films and nanotechnologies [25–27], and as a laser medium [28]. Concerning antimony and its ionization stages, for Stark broadening in literature exist only experimental results for Sb III [29] and theoretical data for Sb II [30].

According to Cohen [31], the cosmic abundance of tellurium (Te) is larger than for any element with atomic number greater than 40, and its spectral lines are identified in stellar spectra. Yuschenko and Gopka [32] found tellurium line in the Procyon photosphere spectrum, Chayer et al. [3] identified tellurium lines in the cool DO white dwarfs HD199499 and HZ21, and Yuschenko et al. [2] detected tellurium in Am star ρ Pup. We note that Stark broadening is important for Am stars and in particular for DO white dwarfs which have effective temperatures from approximately 45,000 K up to around 120,000 K [33]. So that Stark broadening parameters for Te in various ionization degrees are useful. It should be noted as well that tellurium is also interesting as a laser medium [28]. There is no experimental results for Stark broadening of Te spectral lines but exist a study [34] with Stark broadening parameters for Te I multiplets.

In this work we will calculate full widths at half intensity maximum (FWHM), due to collisions with surrounding electrons, for Se IV, Sn IV, Sb IV and Te IV spectral lines, using the modified semiempirical method (MSE) [35–37] and will compare the obtained results with existing theoretical data.

2. The Modified Semiempirical Method

In accordance with the modified semiempirical (MSE) approach [35], the electron impact full width (FHWM) of an isolated ion line is given by the following equation:

$$w_{MSE} = N \frac{4\pi}{3c} \frac{\hbar^2}{m^2} \left(\frac{2m}{\pi kT} \right)^{1/2} \frac{\lambda^2}{\sqrt{3}} \cdot \left\{ \sum_{\ell_i \pm 1} \sum_{L_{i'} J_{i'}} \vec{\Re}^2_{\ell_i,\ell_i \pm 1} \widetilde{g}(x_{\ell_i,\ell_i \pm 1}) + \sum_{\ell_f \pm 1} \sum_{L_{f'} J_{f'}} \vec{\Re}^2_{\ell_f,\ell_f \pm 1} \widetilde{g}(x_{\ell_f,\ell_f \pm 1}) + \left(\sum_{i'} \vec{\Re}^2_{ii'} \right)_{\Delta n \neq 0} g(x_{n_i,n_i + 1}) + \left(\sum_{f'} \vec{\Re}^2_{ff'} \right)_{\Delta n \neq 0} g(x_{n_f,n_f + 1}) \right\}.$$
(1)

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Here, *i* and *f* are for initial and final levels, respectively, $\vec{\Re}^2_{\ell_k,\ell_{k'}}$, k = i, f is the square of the matrix element, and

$$\left(\sum_{k'} \vec{\Re}_{kk'}^2\right)_{\Delta n \neq 0} = \left(\frac{3n_k^*}{2Z}\right)^2 \frac{1}{9} (n_k^{*2} + 3\ell_k^2 + 3\ell_k + 11)$$
(2)

(in Coulomb approximation).

In Equation (1)

$$x_{l_k,l_{k'}} = \frac{E}{\Delta E_{l_k,l_{k'}}}, k = i, f$$

where $E = \frac{3}{2}kT$ is the electron kinetic energy and $\Delta E_{l_k,l_{k'}} = |E_{l_k} - E_{l_{k'}}|$ is the energy difference between levels l_k and $l_k \pm 1$ (k = i, f),

$$x_{n_k,n_k+1} \approx \frac{E}{\Delta E_{n_k,n_k+1}}$$

where for $\Delta n \neq 0$ the energy difference between energy leves with n_k and n_{k+1} , $\Delta E_{n_k,n_k+1}$, is estimated as $\Delta E_{n_k,n_k+1} \approx 2Z^2 E_H/n_k^{*3}$. With $n_k^* = [E_H Z^2/(E_{ion} - E_k)]^{1/2}$ is denoted the effective principal quantum number, Z is the residual ionic charge, for example Z = 1 for neutral atoms, E_{ion} is the appropriate spectral series limit, N and T are electron density and temperature, while g(x) [22] and $\tilde{g}(x)$ [35] are the corresponding Gaunt factors.

3. Results and Discussion

All atomic energy levels needed for calculation of Se IV Stark line widths were found in Rao and Badami [38] and the newest value of ionization potential in Joshi and George [39]; for Sn IV, Sb IV and Te IV energy levels are taken from Moore [40], but the newest values for ionization potential have been taken for Sb IV [41] and for Te IV [42] as suggested in NIST database [43]. The needed matrix elements have been calculated within Coulomb approximation [44].

For the calculation of line width the modified semiempirical method [35] (see also the review of inovations and applications in [37]) has been used. The obtained results for 2 Se IV, 6 Sn IV, 2 Sb IV and 1 Te IV transitions are shown in Table 1 for perturber density of 10^{17} cm⁻³ and temperatures from 10,000 K up to 160,000 K.

There is no experimental or theoretical data for comparison for Se IV, Sb IV and Te IV. For Sn IV exist four transitions where the comparison with semiempirical data of de Andrés-García et al. [21] is possible, while for other two there is no theoretical or experimental data in literature. There are two reasons to calculate Stark broadening for four tansitions already calculated by de Andrés-García et al. [21]. The first step is to compare with their results and to see if a systematic difference exist, so that user who needs also data for two transitions not calculated by [21], could eventualy perform the corresponding scaling or correction. The second reason is to check large difference between Stark widths within the multiplet $5p^2P^0-6s^2S$. For four transitions, existing in the paper of de Andrés-García et al. [21] we performed calculations for temperatures given in this article and compared the corresponding FWHM in Table 2. One can see that differences are up to factor of two and that they decrease towards higher temperatures. Since tin is a complex element and both methods are approximative in comparison with semiclassical perturbation method (see e.g., [45–47] these differences are more or less acceptable and obviously, more experimental date are needed in order to check and improve the theory, which works better for simpler spectra. The agreement is best for Sn IV $6s^2S_{1/2}-6p^2P_{1/2}^0$ line.

Element	Transition	λ (Å)	T(K) = 10,000	20,000	40,000	80,000	160,000
			FWHM(Å)				
Se IV	4p ² P ^o -4d ² D	654.8	0.224×10^{-2}	0.159×10^{-2}	0.112×10^{-2}	0.794×10^{-3}	0.600×10^{-3}
Se IV	5s ² S-5p ² P ^o	2987.7	0.230	0.163	0.115	0.831×10^{-1}	0.697×10^{-1}
Sn IV	$5s^2S_{1/2}-6p^2P_{1/2}^0$	505.4	0.581×10^{-2}	0.411×10^{-2}	0.290×10^{-2}	0.215×10^{-2}	0.182×10^{-2}
Sn IV	$5s^2S_{1/2}-6p^2P_{3/2}^0$	499.9	0.592×10^{-2}	$0.419 imes 10^{-2}$	0.296×10^{-2}	0.218×10^{-2}	$0.183 imes 10^{-2}$
Sn IV	$6s^2S_{1/2}-6p^2P_{1/2}^{o}$	4217.3	0.687	0.486	0.344	0.263	0.229
Sn IV	$6s^2S_{1/2}-6p^2P_{3/2}^{o}$	3862.2	0.591	0.418	0.295	0.225	0.196
Sn IV	$5p^2P_{1/2}^0-6s^2S_{1/2}$	956.3	0.142×10^{-1}	0.101×10^{-1}	0.712×10^{-2}	0.541×10^{-2}	0.460×10^{-2}
Sn IV	$5p^2P_{3/2}^{o}-6s^2S_{1/2}$	1019.7	0.167×10^{-1}	0.118×10^{-1}	0.834×10^{-2}	0.633×10^{-2}	0.536×10^{-2}
Sb IV	5s ¹ S-5p ¹ P ^o	1042.2	0.623×10^{-2}	0.441×10^{-2}	0.312×10^{-2}	0.220×10^{-2}	$0.167 imes 10^{-2}$
Sb IV	6s ³ S-6p ³ P ^o	3543.2	0.429	0.303	0.214	0.158	0.135
Te IV	6s ² S-6p ² P ^o	3375.0	0.366	0.259	0.183	0.134	0.114

Table 1. FWHM - Full Width at Half intensity Maximum (Å) for Se IV, Sn IV, Sb IV and Te IV spectral lines, for a perturber density of 10^{17} cm⁻³ and temperatures from 10,000 to 160,000 K. Calculated wavelength (λ) of the transitions (in Å) is also given.

Table 2. Comparison between FWHM - Full Width at Half intensity Maximum (Å) for Sn IV calculated in this work (W_{TW}) and in de Andrés-García et al. [21] (W_{AAC}), for a perturber density of 10^{17} cm⁻³.

Element	Transition	T(K)	$W_{TW}(\text{\AA})$	$W_{AAC}(A)$
Sn IV	$5s^2S_{1/2}-6p^2P_{1/2}^0$	11,000	0.554×10^{-2}	0.121×10^{-1}
	- 1/2	17,500	0.439×10^{-2}	0.880×10^{-2}
	$\lambda = 505.4$ Å	20,000	0.411×10^{-2}	0.810×10^{-2}
		30,000	0.335×10^{-2}	0.620×10^{-2}
		50,000	0.261×10^{-2}	0.450×10^{-2}
Sn IV	$6s^2S_{1/2}-6p^2P_{1/2}^0$	11,000	0.655	0.888
	- 1/2	17,500	0.520	0.656
	$\lambda = 4217.3$ Å	20,000	0.486	0.603
		30,000	0.397	0.469
		50,000	0.310	0.348
Sn IV	$5p^2P^o_{1/2}-6s^2S_{1/2}$	11,000	$0.136 imes 10^{-1}$	$0.290 imes 10^{-1}$
		17,500	$0.108 imes 10^{-1}$	$0.213 imes 10^{-1}$
	$\lambda = 956.3$ Å	20,000	$0.101 imes 10^{-1}$	$0.195 imes 10^{-1}$
		30,000	0.822×10^{-2}	0.151×10^{-1}
		50,000	0.642×10^{-2}	$0.111 imes 10^{-1}$
Sn IV	$5p^2P^0_{3/2}-6s^2S_{1/2}$	11,000	$0.159 imes 10^{-1}$	$0.433 imes 10^{-1}$
	- 0/2	17,500	0.126×10^{-1}	$0.317 imes 10^{-1}$
	$\lambda = 1019.7 \text{ \AA}$	20,000	0.118×10^{-1}	$0.290 imes 10^{-1}$
		30,000	0.963×10^{-2}	0.223×10^{-1}
		50,000	0.752×10^{-2}	$0.163\times\!10^{-1}$

Concerning the big difference of Stark widths within some multiplets of de Andrés-García et al. [21], we have data for three Sn IV multiplets enabling to check how similar are Stark broadening parameters. Wiese and Konjević [48] concluded in their article where regularities and similarities in plasma broadened spectral line widths have been considered that "line widths in multiplets usually agree within a few per cent". In order to see how it is in the case of Sn IV for multiplets considered in this work. First of all we should convert results from Å units to angular frequency units. For this purpose the following formula can be used:

$$W(\mathring{A}) = \frac{\lambda^2}{2\pi c} W(s^{-1})$$
⁽²⁾

where c is the speed of light.

For the multiplet $5s^2S-6p^2P^{\circ}$, for T = 20,000 K, we obtain in units $10^{12} s^{-1} 0.303$ for $\lambda = 505.4$ Å and 0.316 for $\lambda = 499.9$ Å, so that the difference is 4.1%. For the multiplet $6s^2S-6p^2P^{\circ}$, the corresponding Stark width for $\lambda = 4217.3$ Å is 0.515 and for $\lambda = 3862.2$ Å 0.529, so that the difference is 2.6%. For the multiplet $5p^2P^{\circ}-6s^2S$, the result for $\lambda = 956.3$ Å is 0.208 and for $\lambda = 1019.7$ Å, 0.214 i.e., the difference is 2.8%, which is in accordance with the conclusion of Wiese and Konjević [48]. Contrary, the corresponding values in de Andrés-García et al. [21] are 0.402 and 0.526, respectively, so that the difference is 24%.

We note also that Se IV and Te IV are homologous ions and if we look at Stark widths expressed in angular frequency units for homologous transitions, $5s^2S-5p^2P^0$ for Se IV and $6s^2S-6p^2P^0$ for Te IV widths are only for 11% smaller for Se IV at 10,000 K, and for 30% at 160,000K. Consequently for a rough estimation of Stark width, we could use the value for corresponding transition in homologous atom or ion, nearest to the considered one.

It is worth to add that the theoretical resolving power of the high-resolution echelle spectrometer for the Keck Ten—Meter Telescope is of the order of > 250,000. However, practical realizations may be approximately 36,000. Resolution needed for Stark widths shown in Table 1 may be divided in two groups. For lines between 4217.3 Å and 2987.7 Å needed resolutions are 6,139–12,990 for 10,000 K and 16,035–35,953 for 80,000 K. For lines between 1042.2 Å and 505.4 Å needed resolutions are 61,060–292,321 for 10,000 K and 161,090–824,685 for 80,000 K. It is important that for Sn IV 4217.3 Å line, which is in the visible, needed resolution is 6,139 at 10,000 K and 16,035 at 80,000 K, so that the influence of Stark broadening can be observed with large terrestrial telescopes. For this line, thermal Doppler width is 0.0277 Å and Stark 0.687 Å for T = 10,000 K. The corresponding values for 80,000 K are 0.0784 Å and 0.263 Å respectively.

The obtained Se IV, Sn IV, Sb IV and Te IV Stark widths presented in Table 1, will be implemented as well in the STARK-B database [49,50], which is first of all dedicated for the investigations, modelling and diagnostics of the plasma of stellar atmospheres, but this collection of Stark broadening parameters is also useful for diagnostics of laboratory plasmas, as well as for investigation of laser produced, inertial fusion plasma and for plasma technologies.

We note that STARK-B database participates in the Virtual Atomic and Molecular Data Center— VAMDC [51,52], which enables a more effective search and mining of atomic and molecular data from different databases.

Stark line widths calculated in this work contribute also to the creation of a set of such data for the largest possible number of spectral lines, since this is of importance for a number of problems like stellar spectra analysis and synthesis, opacity calculations and modelling of stellar atmospheres.

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Article Stark Widths of Na IV Spectral Lines

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Abstract: Sodium is a very important element for the research and analysis of astrophysical, laboratory, and technological plasmas, but neither theoretical nor experimental data on Stark broadening of Na IV spectral lines are present in the literature. Using the modified semiempirical method of Dimitrijević and Konjević, here Stark widths have been calculated for nine Na IV transitions. Na IV belongs to the oxygen isoelectronic sequence, and we have calculated Stark widths belonging to singlets, triplets, and quintuplets, as well as with different parent terms. This is used to discuss similarities within one transition array with different multiplicities and parent terms. Additionally, calculated widths will be implemented in the STARK-B database (http://stark-b.obspm.fr) which is also a part of the Virtual Atomic and Molecular Data Center (VAMDC—http://www.vamdc.org/).

Keywords: stark broadening; line profiles; atomic data; Na IV

1. Introduction

In spite of the fact that sodium is important for the research and analysis of various astrophysical (e.g., [1]), laboratory (e.g., [2]), and technological (e.g., [3]) plasmas, Stark broadening data for its different ionization stages are very scarce or even missing. Experimental results exist only for Na I [4]. There are also a number of theoretical results for neutral sodium. For example, semiclassical perturbation (SCP) Stark broadening parameters from [5] have been used for non-LTE calculations for neutral Na in late-type stars [6]. For Na II, there are the semiclassical results of Jones et al. [7] published also by Griem [8], while for Na VI, Na IX, and Na X, there are theoretical SCP results in the STARK-B database [9,10]. There are also quantum mechanical calculations of Stark width for one Na VII, and one Na VIII spectral line [11], and that is all. There are no results for Na IV and Na V.

The creation of a set of Stark broadening parameters for as large as possible number of spectral lines is useful for a number of problems such as stellar spectra analysis and synthesis, opacity calculations, and the modelling of stellar atmospheres. In order to contribute to this aim , we will calculate full widths at half intensity maximum (FWHM), due to collisions with surrounding electrons, for nine Na IV spectral lines using the modified semiempirical method (MSE) [12–14], since a set of atomic data needed for an adequate application of the more sophisticated semiclassical perturbation method [15–17] does not exist. The obtained results will be used for a consideration of the regular behavior of Stark widths within a transition array.

2. The Modified Semiempirical Method

For calculation of Na IV Stark widths, we will use the modified semiempirical method (MSE) [12–14]. Since it is analysed in more details in Dimitrijević et al. [18], only basic information will be summarized in this work. The expression for the electron impact full width (FHWM) of an isolated ion line is [12]:

$$w_{MSE} = N \frac{4\pi}{3c} \frac{\hbar^2}{m^2} (\frac{2m}{\pi kT})^{1/2} \frac{\lambda^2}{\sqrt{3}} \times \{ \sum_{\ell_i \pm 1} \sum_{L_{i'} J_{i'}} \tilde{\Re}^2_{\ell_i, \ell_i \pm 1} \tilde{g}(x_{\ell_i, \ell_i \pm 1}) + \sum_{f \pm 1} \sum_{L_{f'} J_{f'}} \tilde{\Re}^2_{\ell_f, \ell_f \pm 1} \tilde{g}(x_{\ell_f, \ell_f \pm 1}) + (\frac{3n_i^*}{2Z})^2 \frac{1}{9} (n_i^{*2} + 3\ell_i^2 + 3\ell_i + 11)g(x_{n_i, n_i + 1}) + (\frac{3n_f^*}{2Z})^2 \frac{1}{9} (n_f^{*2} + 3\ell_f + 11)g(x_{n_f, n_f + 1}) \},$$

$$(1)$$

where *i* and *f* are for initial and final levels, $\vec{\Re}_{\ell_k,\ell_{k'}}^2$, k = i, f is the square of the matrix element, $x_{l_k,l_{k'}}$ and x_{n_k,n_k+1} are the ratios of electron kinetic energy and the energy difference between the corresponding energy levels (see [18]), *N* and *T* are electron density and temperature, and g(x) [19] and $\tilde{g}(x)$ [12] are the corresponding Gaunt factors.

3. Results and Discussion

The required atomic energy levels for Na IV have been taken from Sansonetti [20], and the corresponding matrix elements have been calculated within the Coulomb approximation [21]. The results for the Stark widths of nine transitions from the Na IV spectrum—obtained by using the modified semiempirical method [12] (see also the review of innovations and applications in [14])—are given in Table 1 for perturber density of 10^{17} cm⁻³ and temperatures from 10,000 K up to 160,000 K. These data are the first for the Stark broadening of Na IV, so there are no other experimental or theoretical data to compare with the present values.

Table 1. FWHM (full width at half intensity maximum; Å) for Na IV spectral lines, for a perturber density of 10^{17} cm⁻³ and temperatures from 10,000 to 160,000 K. Calculated wavelength (λ) of the transitions (in Å) is also given.

Element	Transition	λ (Å)	T (K) = 10,000	20,000	40,000	80,000	160,000
			FWHM (Å)				
Na IV	$(^{2}D)3s^{1}D^{0}-(^{2}D)3p^{1}D$	1534.5	0.259E-01	0.183E - 01	0.130E - 01	0.916E-02	0.705E - 02
Na IV	(² D)3s ¹ D ^o -(² D)3p ¹ F	2156.4	0.466E - 01	0.329E - 01	0.233E-01	0.165E - 01	0.126E - 01
Na IV	$(^{2}P)3s^{1}P^{0}-(^{2}P)3p^{1}P$	1998.6	0.408E - 01	0.289E - 01	0.204E - 01	0.144E - 01	0.109E - 01
Na IV	(² P)3s ¹ P ^o -(² P)3p ¹ D	1791.6	0.337E-01	0.238E-01	0.169E - 01	0.119E - 01	0.903E - 02
Na IV	$({}^{4}S^{o})3s^{3}S^{o}-({}^{4}S^{o})3p^{3}P$	2019.3	0.425E - 01	0.300E - 01	0.212E - 01	0.150E - 01	0.114E - 01
Na IV	(² D)3s ³ D ^o -(² D)3p ³ D	2111.7	0.425E - 01	0.300E - 01	0.212E - 01	0.150E - 01	0.114E - 01
Na IV	(² D)3s ³ D ^o -(² D)3p ³ F	1971.2	0.376E - 01	0.266E - 01	0.188E - 01	0.133E-01	0.101E - 01
Na IV	(² P)3s ³ P ^o -(² P)3p ³ D	1985.9	0.382E - 01	0.270E - 01	0.191E - 01	0.135E - 01	0.102E - 01
Na IV	$({}^{4}S^{o})3s^{5}S^{o}-({}^{4}S^{o})3p^{5}P$	1963.6	0.367E-01	0.259E-01	0.183E-01	0.130E-01	0.976E-02

All calculated data belong to one transition array: 3s–3p, but they have not only different multiplicities, but also different parent terms. This is an interesting set of data which can be used to test how similar Stark broadening parameters are. Wiese and Konjević [22] concluded in their article where regularities and similarities in plasma broadened spectral line widths were considered that "line widths within transition arrays normally stay within a range of about $\pm 40\%$ ", while for supermultiplets, Stark line widths are within about 30%. In order to see how it is in the case of the

Na IV 3s–3p transition array considered in this work, first we should convert results from Å units to angular frequency units. For this purpose, the following formula can be used:

$$W(Å) = \frac{\lambda^2}{2\pi c} W(s^{-1})$$
⁽²⁾

where *c* is the speed of light. The results in angular frequency units—convenient for the consideration of regularities—are shown in Table 2. We can see that the lowest value is 13.5% smaller than the largest one at 10,000 K and for 15.4% at 160,000 K. Within the data for the considered transition array, there are three cases when two multiplets belong to the same supermultiplet. Namely, two singlets with the parent term (²D), two singlets with the parent term (²D). For them, the lower value is 8.7%, 3%, and 1.6% smaller from the larger one at 10,000 K and 9.4%, 2.6%, and 1.2% at 160,000 K. This is in excellent accordance with conclusions of Wiese and Konjević [22], and we can conclude that missing values for other lines from the considered transition array are similar, and if needed, averaged data from Table 2 (in angular frequency units) can be used for the estimation of their Stark widths.

Table 2. Same as in Table 1, but FWHM is in angular frequency units.

Element	Transition	λ (Å)	T (K) = 10,000	20,000	40,000	80,000	160,000
			FWHM (s $^{-1}$) $ imes$ 10 $^{-11}$				
Na IV	(² D)3s ¹ D ^o -(² D)3p ¹ D	1534.5	2.07	1.47	1.04	0.733	0.564
Na IV	(² D)3s ¹ D ^o -(² D)3p ¹ F	2156.4	1.89	1.33	0.943	0.667	0.511
Na IV	(² P)3s ¹ P ^o -(² P)3p ¹ P	1998.6	1.92	1.36	0.962	0.681	0.516
Na IV	(² P)3s ¹ P ^o -(² P)3p ¹ D	1791.6	1.98	1.40	0.989	0.700	0.530
Na IV	$({}^{4}S^{o})3s^{3}S^{o}-({}^{4}S^{o})3p^{3}P$	2019.3	1.96	1.39	0.981	0.694	0.526
Na IV	(² D)3s ³ D ^o -(² D)3p ³ D	2111.7	1.79	1.27	0.897	0.634	0.483
Na IV	(² D)3s ³ D ^o -(² D)3p ³ F	1971.2	1.82	1.29	0.911	0.644	0.489
Na IV	(² P)3s ³ P ^o -(² P)3p ³ D	1985.9	1.83	1.29	0.913	0.645	0.487
Na IV	$({}^{4}S^{o})3s^{5}S^{o}-({}^{4}S^{o})3p^{5}P$	1963.6	1.79	1.27	0.895	0.633	0.477

Our values of Stark widths for Na IV spectral lines will be added to the STARK-B database [9,10], which is also a part of the set of databases included in the Virtual Atomic and Molecular Data Center (VAMDC) [23,24], enabling a much better and easier search and mining of atomic and molecular data.

We hope that the new Stark broadening data obtained in this work will be of interest for a number of problems in astrophysics and for the diagnostics of laboratory plasmas, as well as for the investigation of laser-produced inertial fusion plasma and for plasma in industry (e.g., laser welding or plasma-based light sources).

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Article Regularities and Systematic Trends on Zr IV Stark Widths

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Abstract: Regularities and systematic trends among the Stark widths of 18 Zr IV spectral lines obtained by modified semiempirical approach have been discussed. Also we compared those calculated Stark broadening parameters with estimates according to Cowley, Purić et al. and Purić and Šćepanović and checked the possibility to find some new estimates. It is demonstrated as well that the formula of Cowley (1971) overestimates Stark widths, obtained by using modified semiempirical method, with the increase of angular orbital momentum quantum number due to its neglection. It is also found that the results obtained by using formula for simple estimates of Purić et al. (1991) are in agreement with the modified semiempirical results within the estimated error bars of both methods, while the estimates using formula of Purić and Šćepanović (1999) are in strong disagreement which increases of angular orbital momentum quantum number.

Keywords: Stark broadening; line profiles; atomic data

1. Introduction

Stark broadening theory is very important for many laboratory and astrophysical plasma investigations, since the collisional processes between the charged particles contribute significantly to the spectral line broadening in high-temperature dense plasma. However, available numerous data on Stark widths of spectral lines are up to now still incomplete and despite of a great effort from different groups including French and Belgrade Schools and their co-workers (see e.g., [1–3] and references therein) in determination of Stark widths and shifts for spectral lines of many atoms and ions, the further work on the determination of missing Stark broadening data for applications in astrophysics and plasma physics is still needed.

When the required Stark broadening parameters are missing, and there is no data for the adequate use of more accurate theoretical methods, simple estimates seem very useful. Such quick estimates of Stark-broadening parameters may be based on different regularities and systematic trends among the Stark widths of atomic spectral lines (see e.g., [4]). We focus on present work on searching regularities and systematic trends of Stark widths for 18 Zr IV spectral lines calculated in [5] by modified semiempirical approach (MSE—Dimitrijević & Konjević [6]). This study has two directions. In the first, we compare existing calculated MSE results with estimates made in this work by formulae of Cowley [7], Purić et al. [8] and Purić & Šćepanović [9]. In the second, we try to find estimates based on systematic trends among 18 calculated MSE results. We note as well that there is no experimental Stark broadening data in the literature.

2. Methods

In all of our estimates in this work, the condition that temperatures are near threshold for collisional excitation or below, when Gaunt factor is constant, is satisfied.

$$3kT/2|E_j - E_{j'}| < 2, (1)$$

where *k* is Boltzmann constant and j' is the perturbing level closest to initial or final level. In such a case, from MSE and the semiempirical method follows (see for further details Dimitrijević & Konjević [6] and Griem [10]) that the dependence of Stark broadening parameters with temperature is

$$f(T) = T^{-1/2}.$$
 (2)

For Zr IV lines considered in this work, the condition (1) is in most cases satisfied up to around 20,000 K, for some cases even 30,000 K, so that for all our analysed widths calculated for temperature T = 10,000 K, the condition (1) is satisfied.

Estimation methods used in present research can be divided into two groups: (a) estimates derived from theory (b) estimates based on purely statistical analysis of existing data. We search the estimation formula in two versions. First, in dependence on effective ionization potential:

$$W_{E_1} = a_1 Z^{c_1} \lambda^2 N f(T) (E_{ion} - E_j)^{-b_1}$$
(3)

Second, in dependence on effective quantum number:

$$W_{E_2} = a_2 Z^{c_2} \lambda^2 N f(T) (n_i^*)^{-b_2}.$$
(4)

The relation between effective principal quantum number and upper or lower state is given by:

$$n_i^{*2} = Z^2 [E_H / (E_{ion} - E_j)],$$
 (5)

where Z - 1 is ionic charge, n^* —effective principal quantum number, W_E estimated Stark width in Å, λ wavelength in Å, N-perturber density in m⁻³, T temperature in K, E_H —hydrogen atom energy, E_{ion} —ionization energy, E_j —energy of upper (j = upper) and lower (j = lower) level. Coefficients a, b and c in Equations (3) and (4) are independent of temperature, ionization potential and electron density for a given transition.

First we will estimate Stark widths with the simple formula of Cowley [7] which is often in use by some modern authors in original or modified form (see, for example, Killian et al. [11], Ziegler et al. [12] and Przybilla et al. [13]) and which has recently used by our team for comparison with MSE calculated values of Stark widths of Lu III spectral lines too, see Majlinger et al. [14]. In Majlinger et al. [5] we derived it from the Griem [10] semiempirical formula at low temperature limit and added the contribution to the width of the lower energy level. In such a case, the difference between original formula of Cowley and the modified formula in [5] is for a factor $2/3^{3/2}$ plus the contribution of the lower energy level. Here, we want to test the original formula of Cowley. After rearranging the original equation of Cowley [7] to express width in Å, we obtain:

$$W_{\text{Cow}} = \frac{h^2}{2\pi c} (1/2m^3\pi k)^{1/2} \frac{\lambda^2 N}{Z^2\sqrt{T}} (n_i^{*4})$$
(6)

where *h* is Planck's constant, *c* speed of light and m mass of perturber.

The Cowley's formula is, as we can see, of the type of Equation (4), where j = upper level, and parameters are $a_2 = 2.92 \times 10^{-30}$, $b_2 = 4$ and $c_2 = -2$.

Further, we test the possibility to use the statistical estimates of [8,9] for quick prediction of unknown Stark widths. Comparing the great amount of Stark width data from STARK-B database [1,2], Purić and his co-workers found the correlation between Stark width and difference between ionization energy and energy of the upper state (which is, according to Purić, called upper effective ionization potential) and offered a set of different estimation formulae. Thus we compared our results from MSE calculations with two different approximations: so-called "generalized" estimate (Purić & Šćepanović [9], in the rest of this text PS99) and estimate based on statistical regression analysis of multiply charged heavy ions (Purić et al. [8], in the rest of this text P91). All of Purić's formulae are of type of Equation (1). Purić's works are shown to be useful in some analysis and comparisons, and investigations of systematic trends and regularities, for example in the cases of searching the regularities and trends via specific homologous sequences [15,16].

The equations for Stark width obtained from Purić's regression analysis of existing set of Stark broadening data was adapted for our purpose. After conversion of width from angular frequency units to Angstroms (see for example Hamdi [17]), coefficient values are recalculated from PS99 and P91, where these coefficients are given for Stark widths measured in s^{-1} . Estimation formulae from PS99 and P91 finally is:

$$W_{PS99} = 3.27 \times 10^{-28} Z^{5/2} \lambda^2 N \sqrt{T} (E_{ion} - E_i)^{-3.1}$$
(7)

and

$$W_{P91} = 2.52 \times 10^{-27} Z^0 \lambda^2 N \sqrt{T} (E_{ion} - E_i)^{-1.73}$$
(8)

respectively, if perturber density N is expressed in m⁻³ and energies in eV.

3. Results and Discussion

In Tables 1 and 2, calculated Stark widths of 18 Zr IV lines made by different estimation methods mentioned above are compared to previous MSE calculations of Stark FWHM Majlinger et al. [5]. All results presented in Tables 1 and 2 are calculated for a temperature of 10,000 K and electron density of 10^{23} m⁻³. First we can see that widths W_{PS99} estimated according to Equation (7) are systematically higher than W_{COW} done by Equation (6).

Second, we can see that W_{PS99} does not fit very well to WMSE widths, although this estimate should be generally applicable, the reasons for which we have already explained in detail in Majlinger et al. [5]. On the contrary, much better agreement is found between all of MSE results and results of estimate for multiple charged heavy ions obtained by Equation (8) in P91 (Equation (8) also in this paper), but only if dependence of estimate on residual charge *Z* is neglected (e.g., parameters should be adopted to be c = 0 or Z = 1). In that case, the corresponding ratios of this comparison range from 0.4 to 2.2.

In Table 3, we are presented ranges of values of ratios of W_{COW} , W_{PS99} and W_{P91} with W_{MSE} for different types of transitions. As we can see, ratios between W_{COW} and W_{MSE} range from 0.8 to 4, and they increase with the increase of the orbital angular momentum quantum number ℓ , which is neglected in the Cowley's formula. For s-p transitions differences are from 0.82 up to 2.82, for p-d from 1.57 up to 1.86 and for d-f from 3.74 up to 3.97. Consequently, obtained results demonstrate that for higher ℓ Cowley's formula overestimates the MSE values due to the neglection of ℓ .

For the estimates W_{PS99} the considered ratio for transitions of s-p type ranges from 1.40 to 8.34 while for other considered types of transitions this ratio is from 4.43 up to 24.27. The ratio of W_{P91} with W_{MSE} vary from 0.48 up to 2.13 which is acceptable for a rough estimate.

In articles of Cowley [7] and Purić & Śćepanović [9] there is no discussion of the corresponding error bars. The assumed accuracy of MSE method is within the limits of 50% for simpler spectra while for complex spectra like for Zr IV might be worse. In Purić et al. [8] is stated that the agreement is within 40% error bars for the majority of cases. Consequently the ratio of W_{P91} with W_{MSE} is within mutual error bars.

Transition	λ (Å)	W _{COW} (Å)	W _{COW} /W _{MSE}
5s ² S _{1/2} -5p ² P ^o _{1/2}	2287.38	0.0768	0.91
$5s^{2}S_{1/2}$ - $5p^{2}P^{0}_{3/2}$	2164.36	0.0706	0.92
5s ² S _{1/2} - 6p ² P ^o _{1/2}	760.16	0.0279	2.82
5s ² S _{1/2} -6p ² P ^o _{3/2}	754.39	0.0280	2.75
5p ² P ^o _{1/2} -5d ² D _{3/2}	1546.17	0.0784	1.86
5p ² P ^o _{3/2} -5d ² D _{3/2}	1607.95	0.0847	1.81
5p ² P ^o _{3/2} -5d ² D _{5/2}	1598.95	0.0842	1.82
5d ² D _{3/2} -5f ² F ^o _{5/2}	1836.10	0.324	3.97
5d ² D _{5/2} -5f ² F ^o _{5/2}	1848.03	0.328	3.97
5d ² D _{5/2} -5f ² F ^o _{7/2}	1846.37	0.328	3.97
6s ² S _{1/2} -6p ² P ^o _{1/2}	5781.45	1.62	0.82
6s ² S _{1/2} -6p ² P ^o _{3/2}	5463.85	1.47	0.83
6p ² P ^o _{1/2} -6d ² D _{3/2}	3577.13	1.13	1.61
6p ² P ^o _{3/2} -6d ² D _{3/2}	3795.07	1.27	1.57
6p ² P ^o _{3/2} -6d ² D _{5/2}	3687.95	1.20	1.59
6d ² D _{3/2} -6f ² F ^o _{5/2}	3751.67	2.80	3.74
6d ² D _{5/2} -6f ² F ^o _{5/2}	3775.08	2.83	3.74
6d ² D _{5/2} -6f ² F ^o _{7/2}	3765.39	2.82	3.77

Table 1. Estimated Stark widths for 18 spectral lines of Zr IV according to Cowley (1971) using Equation (6) (W_{COW}) for a temperature of T = 10,000 K and electron density of N = 10^{23} m⁻³. Estimated Stark widths are compared with MSE calculated Stark widths W_{MSE} [5].

Table 2. Estimated Stark widths for 18 spectral lines of Zr IV according to Purić and Šćepanović, 1999 (W_{PS99}) and Purić et al., 1991 (W_{P91}) for a temperature of T = 10,000 K and electron density of N = 10^{23} m⁻³. Estimated Stark widths are compared with MSE calculated Stark widths W_{MSE} [5].

Transition	λ (Å)	W _{PS99} (Å)	W _{P91} (Å)	W _{PS99} /W _{MSE}	W _{P91} /W _{MSE}
5s ² S _{1/2} -5p ² P ^o _{1/2}	2287.38	0.118	0.0530	1.40	0.63
5s ² S _{1/2} -5p ² P ^o _{3/2}	2164.36	0.110	0.0530	1.40	0.63
5s ² S _{1/2} -6p ² P ^o _{1/2}	760.16	0.0826	0.0164	8.34	1.66
5s ² S _{1/2} -6p ² P ^o _{3/2}	754.39	0.0837	0.0164	8.21	1.61
5p ² P ^o _{1/2} –5d ² D _{3/2}	1546.17	0.187	0.0485	4.43	1.15
5p ² P ^o _{3/2} -5d ² D _{3/2}	1607.95	0.202	0.0525	4.31	1.12
5p ² P ^o _{3/2} -5d ² D _{5/2}	1598.95	0.201	0.0521	4.34	1.13
5d ² D _{3/2} -5f ² F ^o _{5/2}	1836.10	1.39	0.173	17.03	2.13
5d ² D _{5/2} -5f ² F ^o _{5/2}	1848.03	1.41	0.176	17.05	2.12
5d ² D _{5/2} –5f ² F ^o _{7/2}	1846.37	1.41	0.176	17.09	2.13
6s ² S _{1/2} -6p ² P _{1/2} ^o	5781.45	4.77	0.950	2.42	0.48
6s ² S _{1/2} –6p ² P ^o _{3/2}	5463.85	4.39	0.862	2.47	0.48
6p ² P ^o _{1/2} -6d ² D _{3/2}	3577.13	4.64	0.611	6.61	0.87
6p ² P ^o _{3/2} -6d ² D _{3/2}	3795.07	5.22	0.688	6.44	0.85
6p ² P ^o _{3/2} –6d ² D _{5/2}	3687.95	4.96	0.652	6.56	0.86
6d ² D _{3/2} -6f ² F ^o _{5/2}	3751.67	18.0	1.36	24.10	1.82
6d 2D5/2-6f 2Fo 5/2	3775.08	18.2	1.37	24.04	1.82
6d 2D5/2-6f 2Fo 7/2	3765.39	18.2	1.37	24.27	1.83

Additionally, following Purić's approach, we tried without success to represent our MSE widths for Zr IV, in angular frequency units, as a polynomial of upper effective principal quantum number or to obtain a linear dependence between them in log-log scale with correlation coefficient above 90% (which proves that the dependence is reliable). However, the correlation coefficient for log-log regression with lower n^* is around 88%:

$$W_{E3} = 1.37 \times 10^{-9} \lambda^2 n_{lower}^{* 2.47}$$
(9)

(in this case, W_{MSE} can be estimated as $W_{MSE} = 1.54 \text{ Å} \times W_{E3} - 0.14$ with correlation coefficient of around 93%) against, for example, around 60% for upper n^* . But we found the polynomial dependence between Stark widths and the corresponding spectral line wavelengths instead, in a form:

$$W_{E4} = C_4 \lambda^4 + C_3 \lambda^3 + C_2 \lambda^2 + C_1 \lambda + C_0$$
(10)

where $C_4 = -0.6 \times 10^{-14}$, $C_3 = 0.63 \times 10^{-10}$, $C_2 = -0.12 \times 10^{-6}$, $C_1 = 0.7 \times 10^{-4}$ and $C_0 = 0.44 \times 10^{-2}$ [5]. Correlation coefficient for this polynomial regression of 4th degree is 99.9%. In all of those estimations parameters are obtained for condition of T = 10,000 K and N = 10^{23} m⁻³.

Unfortunately there is no experimental data for Zr IV. A comparison of our results with reliable experimental data will be of interest so we encourage such measurements.

Table 3. Ranges of values of ratios of $W_{COW},\,W_{PS99}$ and W_{P91} with W_{MSE} for different types of transitions.

Transition	W _{COW} /W _{MSE}	W _{PS99} /W _{MSE}	W _{P91} /W _{MSE}
s–p	0.82-2.82	1.40-8.34	0.48-1.66
p–d	1.57 - 1.86	4.43-6.61	0.85 - 1.15
d–f	3.74-3.97	17.03-24.27	1.82-2.13

4. Conclusions

We tested Cowley's [7] formula and several expressions for quick calculation of electron-impact widths on the basis of regularity and systematic trends, comparing them with MSE results of Zr IV spectral lines calculated elsewhere Majlinger et al. [5]. We demonstrated that Cowley's formula increasingly overestimate Stark width, obtained by using the modified semiempirical method, with the increase of angular momentum quantum number because of its neglection in the Cowley's approach. It is also found that the results obtained by using formula for simple estimates of Purić et al. (1991) are in agreement with the modified semiempirical results within the estimated error bars of both methods, while the estimates using formula of Purić and Šćepanović (1999) are in strong disagreement which increases with the increase of angular orbital momentum quantum number.

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Stark-Zeeman Line Shape Modeling for Magnetic White Dwarf and Tokamak Edge Plasmas: Common Challenges

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Abstract: The shape of atomic spectral lines in plasmas contains information on the plasma parameters, and can be used as a diagnostic tool. Under specific conditions, the plasma located at the edge of tokamaks has parameters similar to those in magnetic white dwarf stellar atmospheres, which suggests that the same line shape models can be used. A problem common to tokamak and magnetic white dwarfs concerns the modeling of Stark broadening of hydrogen lines in the presence of an external magnetic field and the related Zeeman effect. In this work, we focus on a selection of issues relevant to Stark broadening in magnetized hydrogen plasmas. Various line shape models are presented and discussed through applications to ideal cases.

Keywords: line shapes; Stark broadening; Zeeman effect; tokamaks; white dwarfs

1. Introduction

Tokamaks are devices of toroidal shape employed in controlled nuclear fusion research for the confinement of a burning hydrogen plasma, with ion and electron temperatures in the order of 10⁸ K. The ITER project (www.iter.org) aims to demonstrate the principle of producing more energy from the fusion process than is used to initiate it; this large-scale tokamak is presently under construction in France, and it should be operational for first plasma discharge by the end of 2025 [1]. In order to support the operation of the machine, an extensive set of spectroscopic measurements is planned [2]; for example, passive spectroscopy of the Balmer series will provide information on the isotopic proportion of the fueling gas $(N_T/N_D, N_H/N_D)$ in the divertor region [3], where a large amount of neutrals is expected. Under specific conditions (plasma in "detached" regime), the electron density in the divertor can attain values larger than 10^{14} cm⁻³, so that the Stark effect due to the plasma microfield becomes the dominant line-broadening mechanism on lines with a high principal quantum number. Recent observations of D ϵ (transition 7 \rightarrow 2 of deuterium) in the divertor of ASDEX-Upgrade have been used for the determination of the electron density, using the Stark broadening of the line [4]. Balmer lines with a low principal quantum number like $D\alpha$ are also affected by the microfield and Stark broadening enters into competition with both Doppler broadening and Zeeman splitting [5]. The need for accuracy in spectroscopic diagnostics of tokamak edge plasmas has prompted an interest in the development of line broadening models accounting for the simultaneous action of electric and magnetic fields on atomic energy levels, e.g., [6]. Recently, a line shape database for the first Balmer lines

accounting for the Stark and Zeeman effects has been devised for tokamak spectroscopy applications using computer simulations [7]. Similar databases for Stark broadening exist in astrophysics for stellar atmosphere diagnostic applications, e.g., STARK-B (stark-b.obspm.fr) [8]. Magnetic white dwarfs are of particular interest because they present Balmer spectra with Zeeman triplets [9,10] just as in tokamaks, and atmosphere models indicate zones where the electron density value is comparable to that in detached divertor plasmas [11]. Within this context, modeling work by tokamak spectroscopists in conjunction with astrophysicists aimed at improving Stark line shape databases is presently ongoing. In this paper, we report on models and techniques used for the description of Stark-Zeeman line shapes. A special emphasis is put on ion dynamics and collision operator models.

2. Stark Broadening Formalism

The formalism used in Stark line shape modeling involves atomic physics and statistical physics. An atom immersed in a plasma and emitting a photon is considered; during photon emission, the energy levels are perturbed due to the presence of the charged particles surrounding the atom, and this perturbation results in a splitting and broadening of spectral lines. The Doppler effect, which stems from the atom's motion, is not considered, hereafter, for the sake of clarity (it can be accounted for through convolution). According to standard textbooks (e.g., [12]), a line shape $I(\omega)$ is given by the Fourier transform of the atomic dipole autocorrelation function, a quantity described in the framework of quantum mechanics

$$I(\omega) = \frac{1}{\pi} \operatorname{Re} \int_0^\infty dt C(t) e^{i\omega t}$$
(1)

$$C(t) = \sum_{\alpha\alpha\prime\beta\beta\prime\epsilon} \rho_{\alpha\alpha} (\mathbf{d}_{\alpha\beta} \cdot \mathbf{\epsilon}) (\mathbf{d}_{\alpha\prime\beta\prime}^* \cdot \mathbf{\epsilon}) \Big\{ U_{\alpha\prime\alpha}(t) U_{\beta\prime\beta}^*(t) \Big\}$$
(2)

Here, ρ is the restriction of the density operator to the atomic Hilbert space evaluated at initial time, $\mathbf{d} \cdot \mathbf{\epsilon}$ is the dipole projected onto the polarization vector $\mathbf{\epsilon}$, the indices α , α' (resp. β , β') are used for matrix elements and denote states in the upper (resp. lower) level, the brackets { . . . } denote an average over the perturber trajectories, and U(t) is the evolution operator. It obeys the time-dependent Schrödinger equation

$$i\hbar\frac{dU}{dt}(t) = [H_0 + V(t)]U(t)$$
(3)

Here H_0 is the Hamiltonian accounting for the atomic energy level structure and $V(t) = -\mathbf{d}.\mathbf{F}(t)$ is the time-dependent Stark effect term resulting from the action of the microscopic electric field $\mathbf{F}(t)$ and written in the Schrödinger picture, here. When this term is neglected, the Schrödinger equation has the trivial solution $U(t) = \exp(-iH_0t/\hbar)$, which shows, using Equations (2) and (1), that $I(\omega)$ reduces to a set of delta functions (or Lorentzian functions if the natural broadening is retained). By contrast, the case where $\mathbf{F}(t)$ is significant is much trickier, because there is no general exact analytical solution. The time-dependent perturbation theory yields a formal expansion (Dyson series), which is not applicable in explicit calculations because of the non-commutation of the interaction term at different times (time-ordering problem). This concerns, in particular, the microfield due to ions. Several models, based on suitable approximations, have been developed in such a way to provide an analytical expression for the line shape (e.g., the impact and static approximations, the model microfield method, etc.). On the other hand, computer simulations involving the numerical integration of the Schrödinger Equation (3) [13,14] provide reference line profiles and can serve as a benchmark for testing models. In the following, we focus on the ion dynamics issue, which is important for hydrogen lines with a low principal quantum number.

3. Ion Broadening at the Impact Limit

Modeling work has been done over the last decade, in order to set up an analytical formula for the Lyman α and Lyman β line shapes for opacity investigations in tokamak divertor plasmas ([15] and Refs. therein). Kinetic simulations of radiation transport using the EIRENE code (www.eirene.de) have

indicated the possibility of an alteration of the ionization-recombination balance due to the Lyman opacity [16]. The code employs a Monte Carlo method with random walks and related estimators; line shapes in the atomic frame of reference, i.e., without Doppler broadening, are involved in the sampling of photon frequencies. Because of the low principal quantum numbers (n = 2 and 3) involved in Lyman α and Lyman β , the ion dynamics is important, and a collision operator can be used for ions as a first approach for an estimate of the Stark broadening. The model of Griem, Kolb, and Shen (GKS, [17]), developed for electron broadening in hydrogen plasmas, can be adapted in a simple way for ions. Consider a single ion perturbing the emitter (binary interaction); the collision operator *K* reads

$$K = N \int_0^\infty dv f(v) v \int_0^{\lambda_D} db 2\pi b \{1 - S\}_{\text{angle}}$$
(4)

where the integrals are done over the velocity (module) v and the impact parameter (module) b, N and f stand for the perturber's density and velocity distribution function, the brackets denote an angular average and S is the scattering matrix corresponding to a binary collision. The emitter motion is accounted for through a reduced mass model. The upper bound λ_D is the Debye length, which accounts phenomenologically for the screening of the Coulomb field due to the electrons. In the GKS model, the integration domain is restricted in such a way that S can be expanded at the second order in the Stark perturbation $V(t) = -\mathbf{d}.\mathbf{F}(t)$ ("weak collisions"); the integrals can be performed explicitly in the case of a Coulomb field and the resulting collision operator reads (notations are different from those in [17])

$$K_{\text{weak}} = N\pi b_W^2 v_0 \times \frac{2}{\sqrt{\pi}} \frac{r^2}{n^4} E_1 \left[\left(\frac{b_W}{\lambda_D} \right)^2 \times \frac{r^2}{n^4} \right]$$
(5)

Here, $v_0 = \sqrt{(2k_BT/m_{red})}$ is the thermal velocity, accounting for reduced mass; $b_W = (2/3)^{1/2} \hbar n^2/m_e v_0$ is the Weisskopf radius, which discriminates between weak $(b > b_W)$ and strong $(b < b_W)$ collisions (sometimes this is defined without the $(2/3)^{1/2}$ factor); r^2 is the square of the atomic electron position operator, in atomic units, restricted to the subspace relative to quantum number n, and can be formally replaced by its diagonal matrix element $(9/4)n^2(n^2 - l^2 - l - 1)$; and E_1 is the exponential integral function. Equation (5) represents the contribution of weak collisions only; these are usually dominant with respect to strong collisions, that the impact approximation is valid. An estimate of the strong collision contribution to K can be done by setting formally $S \equiv 0$ in Equation (4) (this function strongly oscillates with respect to b and v) and adjusting the integration domain in such a way that only impact parameters smaller than the Weisskopf radius are retained. An explicit calculation yields

$$K_{\text{strong}} = N\pi b_W^2 v_0 \times \frac{2}{\sqrt{\pi}} \frac{r^2}{n^4} \tag{6}$$

and the final expression for the collision operator is $K = K_{\text{weak}} + K_{\text{strong}}$. A Stark line shape described within the impact approximation can be written as a sum of Lorentzian functions with a half-width at half-maximum given by the matrix elements of the collision operator. In the framework of radiative transfer Monte Carlo simulations, this renders the sampling of photon frequencies straightforward. Figure 1 shows a plot of Lyman α at conditions of tokamak edge plasmas, calculated without Doppler broadening and using the GKS collision operator. The result is in contrasted to the result of a computer simulation, where the Schrödinger equation is solved numerically for a given set of microfield histories. These histories are simulated using particles moving in a cube with periodic boundary conditions (see [7] for details on the code). As can be seen, the impact approximation agrees well with the simulation result. Two essential features of the impact model are the assumptions of binary collision and short collision time; these features are characterized by the ratios b_W/r_0 and τ_c/t_i , which must be much smaller than unity. Here, $r_0 \sim N^{-1/3}$ is the mean interparticle distance, $\tau_c \sim r_0/v_0$ is the collision time, and t_i is the time of interest (which is estimated as the inverse line width). Here, the values of

 $3\times10^{12}\,cm^{-3}$ and 1 eV, assumed for density and temperature, respectively, yield values of 4% and 0.6% for these ratios.



Figure 1. Deuterium Lyman α line profile in the atomic frame of reference, i.e., without Doppler broadening, in conditions such that the ratios b_W/r_0 and τ_c/t_i are much smaller than unity (see text). As can be seen, the impact theory agrees well with the simulation.

4. Influence of Zeeman Effect on Line Broadening

In tokamaks, the magnetic field is strong enough that hydrogen lines exhibit a Zeeman triplet structure. This structure is a feature of energy degeneracy removal relative to the magnetic quantum number *m*; it can alter the Stark broadening if the Zeeman perturbation $\hbar\omega_Z = |\mu_B B| (\mu_B \text{ being}$ the Bohr magneton) exceeds the characteristic Stark perturbation, estimated either by $n^2 ea_0 F_0$ (static picture, F_0 being the Holtsmark field) or $\hbar N \pi b_W^2 v_0 \ln(\lambda_D / b_W)$ (impact limit, the logarithm stems from the behavior of the exponential integral for small argument). An extension of the GKS impact collision operator that accounts for this degeneracy removal can be devised through the use of an alternative *S* operator in Equation (4) involving the Zeeman Hamiltonian. The method employs an adaptation of the GBKO model (Griem, Baranger, Kolb, Oertel [18]), initially developed for helium lines, for hydrogen lines with Zeeman effect; details can be found in [6,19]. The result is a collision operator with a dependence on the *m* quantum number. The following relation holds

$$\langle \alpha | K | \alpha \rangle = N \int_0^\infty dv f(v) v \sigma_\alpha(v) \tag{7}$$

$$\sigma_{\alpha} = \pi (b_{\alpha}^{\text{st}})^2 + 2\pi b_{W\alpha}^2 \left\{ K_{\alpha}^{\prime\prime} \ln \left(\frac{\lambda_D}{b_{\alpha}^{\text{st}}} \right) + \left[K_{\alpha}^{\perp} a(s) - i K_{\alpha}^{\prime\perp} b(s) \right]_{s=\lambda_D/b_m}^{s=b_{\alpha}^{\text{st}}/b_m} \right\}$$
(8)

where α stands for the quantum numbers n, l, m; the constants $K_{\alpha}^{//} = \sum_{\alpha'} |z_{\alpha'\alpha}|^2/n_{\alpha}^4$, $K_{\alpha}^{\perp} = \sum_{\alpha'} [|x_{\alpha'\alpha}^+|^2 + |x_{\alpha'\alpha}^-|^2]/n_{\alpha}^4$, and $K'_{\alpha}^{\perp} = \sum_{\alpha'} [|x_{\alpha'\alpha}^+|^2 - |x_{\alpha'\alpha}^-|^2]/n_{\alpha}^4$ involve the atomic electron position operator (here, by convention, $x^{\pm} = (x \pm iy)/\sqrt{2}$); a(s) and b(s) are the GBKO functions writable in terms of the modified Bessel functions, b_{α}^{st} is the strong collision radius; and $b_m = v/\omega_Z$ is a distance relative to Zeeman degeneracy removal. The latter mitigates the characteristic strength of the Stark perturbation. If the ordering $b_{W\alpha} \ll b_m$ is satisfied, the *a* function evaluated at the upper bound can be replaced by a logarithm, and the length b_m plays a role similar to the Debye length. Note that the Weisskopf radius here is a function of v; the definition given in the previous section corresponds to evaluation at the thermal velocity $v = v_0$. In Equation (8), the dependence on velocity has not been written explicitly for the sake of simplicity. The so-called "strong collision radius"—here, b_{α}^{st} —coincides with the Weisskopf radius at the limit $B \rightarrow 0$, and is different at finite magnetic field. It can be determined by the solving of an integral equation; see [6,19] for details. The mitigation of Stark broadening due to the Zeeman effect is all the more important so that the Zeeman energy

sublevels are separated. This effect is particularly important on the lateral components of Lyman α because they are not affected by $\Delta m = 0$ couplings (i.e., $K_{\alpha}^{//}$ vanishes identically in Equation (8)). Figure 2 shows an illustration of the Stark broadening mitigation on the lateral blue component. The same plasma conditions as in the previous section are considered, and a magnetic field of 2 T is assumed. The adaptation of the GBKO model yields a reduction of the line width by a factor of three. A slight shift towards the blue side is also visible; this corresponds to the imaginary part of the collision operator matrix element, and its amplitude is determined by the GBKO *b* function.



Figure 2. Lateral blue component of the Lyman α Zeeman triplet in the atomic frame of reference. A magnetic field of 2 T is assumed and the observation is assumed perpendicular to *B* ($\theta = 90^\circ$). "GBKO" refers to the Griem-Baranger-Kolb-Oertel 1962 model, adapted to hydrogen in the presence of Zeeman effect; "GKS" refers to the original Griem-Kolb-Shen 1959 model for hydrogen without degeneracy removal. As can be seen, the Zeeman degeneracy removal, retained in the GBKO model, strongly reduces the line width. A slight shift is also visible.

5. An Extension of the Impact theory to Non-Binary Interactions

The impact approximation for ions can be inaccurate under high density or low temperature conditions, namely, when the dimensionless parameters $\langle \alpha | K | \alpha \rangle \times \tau_c$ and b_W/r_0 are significant. Comparisons with ab initio simulations have indicated that the use of the impact approximation can result in a systematic overestimate of the Stark width, e.g., [6]. Recently, a model has been developed in such a way to account for the deviations analytically [20]. The model uses kinetic theory techniques and is inspired by the "unified theory", a formalism proposed in the late sixties as an extension of the impact approximation for electrons [21,22]. The main result of the unified theory is a frequency-dependent collision operator, which accounts for incomplete collisions (i.e., collisions not achieved during the time of interest) and provides a correct asymptotic line shape formula. If we apply the unified theory to ions, the collision operator reads

$$K(\omega) = N \int_0^\infty dt e^{i\omega t} \left\{ V(t) e^{-iH_0 t/\hbar} Q(t) V(0) \right\}$$
(9)

where *V* stands for the Stark interaction corresponding to a binary collision with an ion, N is the number of ions, the brackets { . . . } denote statistical average, and *Q*(*t*) is the evolution operator of the atom under the influence of one collision in the interaction picture. It obeys the Schrödinger equation and admits the Dyson series as a formal expression. Several techniques exist for the evaluation of the integral, either based on the exact solution or on the perturbation theory, as done for the impact collision operator. The behavior of the collision operator at small and large frequency detuning is obtained by asymptotic analysis. This shows that *K*(ω) behaves as $|\Delta \omega|^{-1/2}$ in the wings, and reduces to the impact collision operator at the center. For applications in tokamak edge and divertor conditions, it has been necessary to modify the original model Equation (9) in such a way as to account for N body

interaction effects, referred to as "correlated collisions". Such correlations occur if, during one collision, the atom "feels" the presence of the other perturbers. Correlated collisions are important when the characteristic collision frequency $\langle \alpha | K | \alpha \rangle$ becomes of the same order as, or larger than, the inverse correlation time of the emitter-perturber interaction potential [23,24]. The latter is of the order of the inverse plasma frequency ω_{pi}^{-1} (due to Debye screening). In the model reported in [20], correlated collisions are retained through a resummation procedure applied to kinetic equations of BBGKY-type. The resulting collision operator obeys a nonlinear equation with a structure similar to Equation (9)

$$K(\omega) = N \int_0^\infty dt e^{i\omega t} \left\{ V(t) e^{-i(H_0 - iK_0)t/\hbar} Q(t) V(0) \right\}$$
(10)

In the exponential, $K_0 = K(\omega_0)$ is the collision operator evaluated at the central frequency of the line under consideration. Its presence denotes a non-Hermitian part in the atomic Liouvillian, which can be interpreted as a renormalization or "dressing" of the atomic energy levels, induced by the presence of other perturbers during a single collision and their correlation with the collision under consideration. This model presents similarities to the result of the resonance broadening theory used for plasma turbulence [25,26]. In practice, a calculation of the collision operator from Equation (10) should be done by iterations. Tests have indicated that such a procedure yields a fast convergence, with typically no more than ten iterations. Figure 3 presents a plot of Lyman α broadened due to ions at $N_e = 10^{15}$ cm⁻³, $T_e = T_i = 1$ eV, which was obtained using impact approximation, the unified theory Equation (9), and its extension Equation (10) applied to ions. No magnetic field is retained here. A numerical result from a simulation is also shown in the figure. As can be seen, the impact approximation strongly overestimates the width. This stems from the inadequacy of this model as a regime where incomplete and correlated collisions are present. The extended unified theory gives a much better result, with an overestimate of the width no larger than 20%. We can also see that the standard unified theory (i.e., not accounting for correlated collisions) is not sufficient here, and yields a different (and incorrect) shape structure, with a dip at the center. Correlated collisions are important in this region, because they govern the average atomic evolution operator at long times; hence, by virtue of the Fourier transform, at small frequency detuning. The typical range for the dip corresponds to frequencies smaller than the matrix elements of K_0 .



Figure 3. Profile of the Lyman α line at conditions such that neither the impact approximation nor the unified theory (UT) for ions are valid. The extension of the unified theory (UT), which accounts for correlated collisions, provides a good estimate of the line width.

6. Stark-Zeeman Line Shapes in Magnetic White Dwarfs

About 10% of white dwarfs are known to have a magnetic field strength of 10^5 to 5×10^8 G (10 to 5×10^4 T), as indicated by spectroscopic observations and models [9,10,27]. An interpretation of the shape of absorption lines requires the Zeeman effect be accounted for in line broadening

models, as done in tokamak edge plasma spectroscopy. Work is currently ongoing in order to improve already-existing databases by implementing the Zeeman effect in Stark line shape models. Figure 4 shows an illustration of the importance of the Zeeman effect in white dwarf spectra. A profile of H α has been calculated with a numerical simulation assuming $N_e = 10^{17}$ cm⁻³, $T_e = T_i = 0.5$ eV, B = 500 T and observation perpendicular to *B*. These conditions are characteristic of the atmosphere of a white dwarf of DAH (magnetic DA) type, presenting hydrogen absorption lines. The spectrum exhibits a triplet structure characteristic to the Zeeman effect. The description of Stark broadening can be more challenging than in tokamak devices because particle correlations are important (here r_0/λ_D is of the order of unity) and the quasiparticle model used in the simulation may not be accurate enough. An extension of this work would consist of using full molecular dynamic simulations. Features specific to dense plasmas, such as quadrupole interactions, could also be important, and should be addressed in detail; see [28] for recent works. The presence of a strong magnetic field changes the plasma dynamics through the gyromotion effect and this can result in an alteration of electron collision operator formulas [29,30]. Specific modifications to the atomic energy level structure (quadratic Zeeman effect, Lorentz electric field) could also be important [31].



Figure 4. Balmer α line shape in conditions relevant to DAH white dwarf atmospheres. The spectrum exhibits a triplet structure due to the strong magnetic field.

7. Non-Hydrogen Species

Tokamak plasmas contain a sufficient amount of impurities, due to plasma-wall interactions or vacuum chamber conditioning, to be observable on spectra (e.g., oxygen, nitrogen, carbon [32]). Specific atomic species of high Z can be found in some, but not all, tokamaks (e.g., tungsten [33]). Helium is also expected to be present in high amounts in ITER during the nuclear phase, where discharges with tritium will be done. As a rule, an analysis of the position and the intensity of impurity lines provides information on the plasma composition, the recycling and sputtering processes, which helps in the setting up of tokamak discharge scenarios. The design of fast routines employs line shape codes (e.g., PPP [34]), and can also use databases available in astrophysics such as STARK-B (stark-b.obspm.fr). An issue that is specific to non-hydrogen lines concerns the modeling of collisional electron broadening, accounting for degeneracy removal with quantum number l. This degeneracy removal yields a reduction of the Stark effect and can induce a shift on the line under consideration. An illustration of this effect is shown in Figure 5. The He 492 nm line (1s4d ¹D–1s2p ¹P) has been addressed, assuming $N_e = 10^{15}$ cm⁻³ and $T_e = 10,000$ K, without ion broadening and without magnetic field for the sake of clarity. Two Lorentzian curves are displayed in the graph. One corresponds to the result of a hydrogen approximation (through the GKS impact model discussed above), and the other one corresponds to the model reported in the STARK-B database, which accounts for the degeneracy removal. As can be seen, the line width is sensitive to the model used. The shift visible in the STARK-B

result is a feature of the degeneracy removal. It depends on the plasma parameters, and can also be used for diagnostic purposes. An extension of already-existing databases to magnetized plasma conditions, where the Zeeman effect is visible in spectra, is under consideration. Work is presently ongoing for applications both in magnetic fusion and astrophysical plasmas.



Figure 5. Plot of the electron impact broadening of He 1s4d ¹D–1s2p ¹P calculated using the STARK-B database (solid line) and a hydrogen approximation for the energy level structure (dashed line). Conditions relevant to high-density tokamak divertor plasmas are considered here, and no magnetic field is retained for the sake of clarity. As can be seen, the result significantly depends on the model used.

8. Conclusions

The shape of atomic spectral lines in plasmas contains information on the plasma parameters, and can be used as a diagnostic tool. In this work, we have examined a selection of problems involved in the modeling of Stark broadening for applications in magnetic fusion and in astrophysics. A specific issue concerns the description of ion dynamics on hydrogen lines, which is important if the upper principal quantum number is low as on Balmer α . The use of a collision operator for ions is convenient because the line shape can be described analytically. In this framework, we have shown that the early model by Griem et al. (GBKO) for atoms with nondegenerate levels can be adapted to hydrogen in the presence of the Zeeman effect. It coincides with numerical simulations at the so-called impact limit, i.e., when the microfield fluctuates at a time scale much smaller than the time of interest, and when strong collisions are rare events. We have also shown that deviations in the impact limit due to non-binary interactions can be retained through an adaptation of the so-called unified theory, a formalism based on kinetic equations and related statistical concepts (BBGKY hierarchy). In astrophysics, magnetic white dwarfs present similarities with tokamak edge plasmas because the hydrogen absorption lines in atmospheres can exhibit a Zeeman triplet structure. We have illustrated this point through calculation of an absorption line in conditions relevant to a DAH stellar atmosphere. In this framework, an extension of already-existing Stark databases that is devoted to accounting for the Zeeman effect is presently under preparation. We have also shown that the description of the broadening of non-hydrogen lines is a challenging issue, both in astrophysics and in magnetic fusion. Modeling work is also ongoing in this framework in order to improve the databases.

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Article A New Analysis of Stark and Zeeman Effects on Hydrogen Lines in Magnetized DA White Dwarfs

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Abstract: White dwarfs with magnetic field strengths larger than 10 T are understood to represent more than 10% of the total population of white dwarfs. The presence of such strong magnetic fields is clearly indicated by the Zeeman triplet structure visible on absorption lines. In this work, we discuss the line broadening mechanisms and focus on the sensitivity of hydrogen lines on the magnetic field. We perform new calculations in conditions relevant to magnetized DA stellar atmospheres using models inspired from magnetic fusion plasma spectroscopy. A white dwarf spectrum from the Sloan Digital Sky Survey (SDSS) database is analyzed. An effective temperature is provided by an adjustment of the background radiation with a Planck function, and the magnetic field is inferred from absorption lines presenting a Zeeman triplet structure. An order-of-magnitude estimate for the electron density is also performed from Stark broadening analysis.

Keywords: line shapes; Stark broadening; Zeeman effect; white dwarfs

1. Introduction

White dwarfs—the end products of stellar evolution—occupy a key position in astrophysical theory. Their properties provide clues to the physical processes that take place during the evolutionary stages near the end of stellar lifetimes. With an understanding of white dwarf magnetic fields, one can indicate the formation or the existence of centered or non-centered dipoles of these magnetic fields [1]. However, only 10% of white dwarfs are thought to carry strong magnetic fields. Most of them have so far only been analyzed by visual comparison of the observation with relatively simple models of the radiation transport in a magnetized stellar atmosphere [2,3]. Some of them have been analyzed by using the least squares minimization scheme to find the best fit for magnetic field geometry [1,4]. Studies of white dwarf atmospheres have shown that the majority of white dwarfs have an atmosphere of pure hydrogen as a result of gravitational setting, which removes helium and heavier elements from the atmosphere and moves them towards inner layers [5,6]. These atmospheres can be considered as hydrogen plasmas, which are similar to some created in laboratory. Such white dwarfs are classified as of DA type due to the strong hydrogen absorption lines they present. The electron density in a white dwarf atmosphere is high enough (up to 10¹⁷ cm⁻³, and higher) so that the line shapes are dominated by Stark broadening, and hence can serve as a probe for the electron density N_e . A noticeable feature of magnetized DA white dwarfs is that their absorption lines exhibit a Zeeman triplet structure, which stems from the perturbation of the atomic energy levels due to the magnetic field. The design of a line shape model accounting for both Stark broadening and Zeeman splitting is not straightforward, because these two effects do not act as additive perturbations. Previous investigations in low-density (tokamak edge) magnetized plasma conditions have indicated an alteration of the broadening of each

Zeeman component due to the change of the atomic energy level structure (degeneracy removal) in response to an external magnetic field [7,8]. This issue also concerns magnetized white dwarfs with spectra exhibiting Zeeman splitting. In this work, we report on the current status of the design of a line shape model accounting for Stark and Zeeman effects simultaneously, for applications to white dwarf atmosphere analysis. We perform new Stark–Zeeman line shape calculations. The applicability of spectroscopy as a diagnostic means is illustrated through the analysis of a white dwarf spectrum obtained from the Sloan Digital Sky Survey (SDSS) database.

2. Stark Line Shape Modeling

In a plasma, the energy levels of atomic species (either neutral atoms or multicharged ions) emitting line radiation is perturbed by the microscopic electric field, which results in a broadening of spectral lines. The spectral profile of an atomic line is proportional to the Fourier transform of the dipole autocorrelation function C(t) [9]

$$L(\omega) = \frac{1}{\pi} \operatorname{Re} \int_0^\infty dt C(t) e^{i\omega t},\tag{1}$$

$$C(t) = \sum_{\vec{e}} \langle \langle \vec{d} \cdot \vec{e} | \{ \hat{U}(t) \} | \rho \vec{d} \cdot \vec{e} \rangle \rangle,$$
⁽²⁾

where $\{...\}$ is the average over plasma particles, ρ is the atomic density operator, and $\vec{d} \cdot \vec{e}$ is the projection of the atomic dipole operator into the polarization plane. The double bracket notation $\langle \langle ... | ... \rangle \rangle$ is used for the scalar product in the Liouville space which is formed by the tensor product $\mathcal{H} \otimes \mathcal{H}^*$ between the atomic state space \mathcal{H} and its dual \mathcal{H}^* . The evolution operator $\hat{U}(t)$ obeys the time-dependent Schrödinger equation in the Liouville space

$$i\frac{d\hat{U}(t)}{dt}(t) = [\hat{L}_0 + \hat{V}(t)]\hat{U}(t),$$
(3)

which is also called the Liouville equation. \hat{L}_0 is the energy superoperator and $\hat{V} = -\hat{d} \cdot \vec{E}$ is the interaction superoperator corresponding to the Stark perturbation. The action of a superoperator \hat{A} on an operator X is defined by $\hat{A}X = [A, X]/\hbar$, where A is the related operator and [,] is the commutator. In a plasma, the electric field is caused by the electrons and the ions and the corresponding Stark perturbation decomposes into two parts

$$\hat{V} = \vec{d} \cdot \vec{E} = -\vec{d} \cdot \vec{E}_i - \vec{d} \cdot \vec{E}_e, \tag{4}$$

where the *e* and *i* subscripts correspond to the electrons and ions, respectively. In the framework of the so-called "standard model", the ions are assumed motionless and a constant electric field is used for their contribution, while the electron contribution is described within a collisional picture by a relaxation ("collision") operator \hat{K}_e , that is through the formal substitution $-\hat{d} \cdot \vec{E}_e \rightarrow -i\hat{K}_e$. In this way, the Liouville equation has the exponential solution $\hat{U}(t) = \exp[-i(\hat{L}_0 - \hat{d}.\vec{E}_i - i\hat{K}_e)]t$ and the line shape function is obtained from a matrix inversion

$$L(\omega) = -\frac{1}{\pi} \operatorname{Im} \sum_{\vec{\epsilon}} \int d^3 E_i W(\vec{E}_i) \langle \langle \vec{d} \cdot \vec{\epsilon} | (\omega - \hat{L}_0 + \hat{\vec{d}} \cdot \vec{E}_i + i\hat{K}_e)^{-1} | \rho \vec{d} \cdot \vec{\epsilon} \rangle \rangle$$
(5)

The integral is performed over the ionic electric field \vec{E}_i and involves its probability density function $W(\vec{E}_i)$. This integration can be performed numerically by using a discretization scheme or by Monte Carlo method. The standard model is convenient for numerical applications due to the structure of Equation (5), which can be interpreted as a sum of complex Lorentzian functions after decomposition of the Liouville double brackets onto a basis is done. The static ion approximation holds

at conditions such that the collision time r_0/v_i (r_0 and v_i being the mean interparticle distance and the thermal ion velocity, respectively) is much larger than the characteristic time for dipole decorrelation (also called "time of interest"). Extensions of the standard model accounting for ion dynamics effects can be set up through suitable modifications; e.g., by formally adding a non-Hermitian contribution to L_0 (Model Microfield Method [10,11], Frequency Fluctuation Model [12,13]). The ion dynamics leads to an additional broadening, which can be important especially at low densities, high temperatures, or for lines with a low upper principal quantum number. Simulations involving a numerical integration of the Liouville Equation (3) and not referring to Equation (5) can also be performed, but they become time-consuming if the atomic system is complex [7,14,15].

For hydrogen lines, the electron collision operator can be evaluated using the Griem–Kolb–Shen model [16], which provides an analytical expression for the contribution of weak collisions based on the time-dependent perturbation series. Such weak collisions involve an impact parameter larger than the Weisskopf radius $b_W = \hbar n^2 / m_e v$ (*n* being the principal quantum number and *v* being the velocity), and they give a dominant contribution to the Stark broadening due to electrons. The strong collision contribution is usually estimated using a Lorentz model, assuming complete interruption of the wave train. Extensions to the Griem–Kolb–Shen model, accounting for "static" perturbations due to the electrons (especially present in the line wings), can be devised through the use of kinetic theory techniques (such as the "unified theory" [17–19]) and they lead to a frequency-dependent collision operator. In white dwarf atmospheres, the plasma is partially ionized and there can be a significant additional broadening caused by collisions with neutrals, either due to resonant interactions or to van der Waals interactions. This broadening is also described with a collision operator \hat{K}_0 , and it is added to \hat{K}_e . Simple models can be found in [9].

3. Zeeman Effect in Magnetized White Dwarfs

In the presence of a magnetic field, the atomic Hamiltonian and the corresponding Liouvillian \hat{L}_0 contain an additional term due to the coupling with the field. Observations of white dwarf spectra have indicated that the magnetic field can attain values larger than 100 T, which renders the Zeeman effect visible on spectra. A description of this effect in line shape modeling requires the following substitution be done in the Liouville Equation (3):

$$\hat{L}_0 \to \hat{L}_0 - \hat{\vec{\mu}} \cdot \vec{B},\tag{6}$$

where $\hat{\mu}$ is the magnetic moment expressed in the Liouville space. We do not consider the weak ("anomalous") Zeeman effect that perturbs fine structure lines, because it is negligible in white dwarf spectra due to high density and strong magnetic fields. If the standard model is used, the same substitution as Equation (6) must be done in the line shape formula (5). An illustration of the Zeeman effect in white dwarf atmosphere conditions is shown in Figure 1. The H α line shape has been calculated assuming an electron density of 10^{17} cm⁻³, a temperature of 1 eV, and a magnetic field of 200 T. An observation direction perpendicular to the magnetic field has been assumed. The line shape exhibits a clean Zeeman triplet structure. The splitting is proportional to the magnetic field, and hence can be used to estimate the magnetic field in white dwarfs. In some white dwarfs, the magnetic field can be sufficiently large so that the quadratic Zeeman effect becomes visible in spectra. This effect can be retained through an additional term in the Hamiltonian which is proportional to B^2 . This term stems from the expansion of the quantity ($\vec{p} + e\vec{A}$)² present in the kinetic energy to the second order in the vector potential \vec{A} . Algebraic manipulations lead to the following expression for the quadratic Zeeman effect Hamiltonian:

$$V_{Z2} = \frac{e^2 B^2 r_{\perp}^2}{8m_e},$$
 (7)

where \vec{r}_{\perp} stands for the atomic electron position operator projected onto the plane perpendicular to \vec{B} . A similar relation holds for the corresponding superoperator in the Liouville space. The quadratic

Zeeman effect is illustrated in Figure 2. A calculation of H α at the same plasma conditions as above, assuming a magnetic field of 2000 T, is shown. Note the asymmetry of the spectrum and the presence of more components.



Figure 1. Plot of H α calculated at $N_e = 10^{17}$ cm⁻³, $T_e = T_i = 1$ eV, with a magnetic field of 200 T. An observation direction perpendicular to the magnetic field has been assumed. The spectrum exhibits a clean Zeeman triplet structure.



Figure 2. Some white dwarfs can have a magnetic field strong enough so that the quadratic Zeeman effect becomes significant. The spectrum becomes asymmetric and presents more components. Here, a calculation has been done assuming B = 2000 T.

4. White Dwarf Spectrum Analysis

In this section, we illustrate the applicability of spectroscopy as a diagnostic means for the characterization of white dwarf atmosphere. A large amount of spectra presenting Zeeman splitting are available on the SDSS database (www.sdss.org) [1,4]. Figure 3 shows an example of such a spectrum. We have performed a fitting of the H α and H β absorption lines using an exponential attenuation model (Beer-Lambert formula) assuming a homogeneous atmosphere slab, leaving the product $N_2 \times L$ between absorber density N_2 and thickness L as an adjustable parameter. The temperature that enters the formula (through the electron collision operator and the microfield probability density function) is estimated from adjustment of the background radiation with a Planck function, assuming local thermodynamic equilibrium. Figure 4 shows the fitting result. The effective temperature is around 7000 K. In order to get a reasonable adjustment on each of these lines, we had to assume different values for the electronic density and the N_2L parameter: $N_e = 5 \times 10^{17}$ cm⁻³ and $N_2L = 5 \times 10^{17}$ m⁻² for H α , $N_e = 2 \times 10^{16}$ cm⁻³ and $N_2L = 2 \times 10^{18}$ m⁻² for H β . A value of 130 T for the magnetic was

found in both cases. These results can be used as a first order-of-magnitude estimate. The necessity of using different values for densities stems from inaccuracies intrinsic to the model (e.g., due to the neglect of self-emission), and this indicates that a further refinement is required.



Figure 3. Plot of the Sloan Digital Sky Survey (SDSS) + J085830 + 412635.12 spectrum in the 4500–7000 Å wavelength range. This spectrum corresponds to a magnetized DA white dwarf. The H α and H β lines present a triplet structure, which is characteristic of Zeeman effect.



Figure 4. An analysis of the shape of absorption lines provides information on the plasma parameters. Here, an adjustment of (a) H α and (b) H β using the Stark-Zeeman line shape model is presented. These lines have been analyzed independently assuming a homogeneous slab geometry. The obtained values for N_e and *B* provide a first order-of-magnitude estimate.

5. Conclusions

According to observations, a significant proportion of white dwarfs have a strong magnetic field, sufficiently so that the absorption lines present a Zeeman triplet structure. In this work, we have done new calculations of Stark–Zeeman hydrogen line shapes in preparation for the development of a model. Specific issues such as the quadratic Zeeman effect need to be investigated further. The applicability of spectroscopy as a diagnostic means for magnetized white dwarf atmospheres has been illustrated through adjustments performed on a spectrum from the Sloan Digital Sky Survey (SDSS) database. It is found that line width adjustments provide an order of magnitude for the electron density and the magnetic field in the atmosphere. An extension of the work to the accounting for plasma non-homogeneity is presently ongoing. Comparisons to other white dwarf spectra analyses, such as in [1,4], will be done.

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Article Line Shape Modeling for the Diagnostic of the Electron Density in a Corona Discharge

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Abstract: We present an analysis of spectra observed in a corona discharge designed for the study of dielectrics in electrical engineering. The medium is a gas of helium and the discharge was performed at the vicinity of a tip electrode under high voltage. The shape of helium lines is dominated by the Stark broadening due to the plasma microfield. Using a computer simulation method, we examine the sensitivity of the He 492 nm line shape to the electron density. Our results indicate the possibility of a density diagnostic based on passive spectroscopy. The influence of collisional broadening due to interactions between the emitters and neutrals is discussed.

Keywords: line shapes; stark broadening; neutral broadening; corona discharge

1. Introduction

Corona discharges of helium were performed in an experiment devoted to the investigation of the dielectric properties of insulators in the context of electrical engineering. This setup consisted of a point-plane electrode system placed inside a helium cryostat; a high voltage was applied to the system and a streamer of either positively or negatively charged particles, ions or electrons, was generated and formed a plasma locally. Here, we report the analysis of the helium 492 nm line (singlet state, 1s4d–1s2p transition) observed in discharges at room temperature. This line is strongly sensitive to the electron density due to the Stark broadening generated by the plasma microfield. We show that a diagnostic of the electron density can be performed from an analysis of the line width and its forbidden component induced by the electric field. This work completes previous research on the analysis of H β [1], and some preliminary results were discussed at the fourth Spectral Line Shape in Plasmas code comparison workshop (SLSP4, see http://plasma-gate.weizmann.ac.il/slsp/). It will be followed with subsequent analyses of spectra observed in helium corona discharges performed at low temperatures (a few K) with liquid helium.

2. Presentation of the Experiment

A point-plane electrode system was placed inside a helium cryostat. The point electrode was negatively polarized by a stabilized high voltage DC power supply (Spellman RHSR/20PN60) and the current-voltage characteristics were measured using a Tektronix TDS540 oscilloscope and a Keithley 610C ammeter. The corona discharge in this geometry is axially symmetric and appears as a luminous spherical region (ionization region) localized near the point electrode against the dark background. More details on the experimental setup can be found elsewhere, e.g., [2]. The measured intensity of the radiation was averaged over the exposure time. A liquid N₂ cooled 2D-CCDTKB-UV/AR detector

was located directly in the exit plane of the spectrograph. The noise level of the CCD detector was determined only by the read-out noise, as the dark current of the camera was less than 1 e/pixel/h at 153 K. The wavelength and intensity response of the detection system was calibrated by using low pressure helium and tungsten ribbon lamps. The line broadening due to the instrument response (1200 g/mm grating) was estimated from the helium lines of a helium lamp as $\Delta\lambda_{ins} < 0.10$ nm.

3. An Analysis of the Helium 492 nm Line Observed at 300 K

In concert with previous research [1], we here report an analysis of the helium 492 nm line (singlet state, 1s4d–1s2p transition) which has been observed on spectra. Discharges were carried out at pressures from 1 to 12 bar, with a current of 100 μ A. In order to simplify the interpretation of the spectra, we focus on the lowest pressure values (namely, between 1 and 2 bar) because the line overlaps with nearby molecular bands at higher pressures.

Figure 1 shows a plot of the helium 492 nm line observed at 1, 1.5, and 2 bar. The line width increases with the pressure. The bump on the blue wing denotes a forbidden transition (1s4f-1s2p) induced by the microfield. An analysis of the resonance broadening resulting from atom-atom collisions using a collision operator model [3] indicates that this effect is not the dominant line broadening one. The instrumental broadening is also not dominant. We examined the role of the Stark effect related to the plasma microfield. A computer simulation method [4] was applied to the He 492 nm line at the same pressure values as above. The He⁺ ion microfield evolution was simulated from a quasiparticle model and the line broadening was calculated from a numerical integration of the time-dependent Schrödinger equation. The contribution of the electrons was evaluated using a collision operator. In our calculations, we used the Griem-Baranger-Kolb-Oertel model [5], assuming an electron temperature of 10⁴ K and leaving the electron density as an adjustable parameter. The ion temperature was assumed to be equal to the atomic (300 K) temperature. Our calculations indicate that the Stark broadening was mainly due to the ions. Figure 2 shows an example of the adjustment performed using the simulation method at p = 1 bar. A value of 1.8×10^{15} cm⁻³ was obtained for the electron density. Our calculations at p = 1.5 bar and 2 bar yielded higher values for the electron density, which indicates an increase trend in terms of the pressure. This result is preliminary and will be completed with further analyses.



Figure 1. Plot of the helium 492 nm line observed in a helium corona discharge at room temperature, for different pressure values. The line width increases with the pressure. This trend is also observed on other lines. The bump on the blue wing denotes a forbidden transition induced by the microfield.



Figure 2. The plasma microfield yields an additional broadening, which can serve as a probe of the density. Here, the Stark broadening was evaluated using a computer simulation method [4].

4. Conclusions

We have analyzed spectral profiles of the helium 492 nm line in helium corona discharges by means of a computer simulation method. An application to 1, 1.5, and 2 bar gas pressures indicates that the plasma microfield yields an obvious Stark broadening, which can serve as a probe for the electron density. This result is a preliminary step in ongoing investigations of liquid corona discharges. New experiments, with liquid helium, are planned and will be analyzed using spectroscopic techniques.

Author Contributions: All authors contributed equally to this work.

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Article



Contribution of Lienard-Wiechert Potential to the Electron Broadening of Spectral Lines in Plasmas

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Abstract: Lienard-Wiechert or retarded electric and magnetic fields are produced by moving electric charges with respect to a rest frame. In hot plasmas, such fields may be created by high velocity free electrons. The resulting electric field has a relativistic expression that depends on the ratio of the free electron velocity to the speed of light in vacuum c. In this work, we consider the semi-classical dipole interaction between the emitter ions and the Lienard-Wiechert electric field of the free electrons and compute its contribution to the broadening of the spectral line shape in hot and dense plasmas.

Keywords: Stark effect; electron broadening; Lienard-Wiechert; retarded interaction; relativistic collision operator

1. Introduction

Line profiles and shifts are used to determine plasma parameters, especially in astrophysics where alternative methods (such as interferometry or Thomson scattering) are not possible. Doppler and pressure broadening (Stark broadening) are typically the two dominant mechanisms and we focus on the latter. In a number of hot astrophysical plasmas, electrons may be energetic enough that their thermal energy K_BT (K_B is the Boltzmann constant and T the temperature) can be comparable to the rest mass. For the extreme densities encountered in some astrophysical objects, pressure broadening could dominate; however, for such objects the electrons may become relativistic due to the extreme temperatures and hence it makes sense to check the modifications to the pressure broadening by relativistic effects. Similarly, the laser-produced plasmas may be achieved in both very high densities and very high temperatures: the first leads to the dominance of Stark over Doppler broadening whereas the second leads to relativistic electron velocities. More specifically, plasma spectroscopy is used in a wide range of electron density from 10 particles per cm³ (interstellar space) to 10^{25} particles per cm³ (star interiors, inertial confinement fusion) and for temperatures between 10⁷ K and 10¹⁰ K. More precisely, as the Doppler effect is constant for a fixed temperature (about 10^8 K in our case), we can neglect it for densities higher than 10^{20} cm⁻³. In the present work, we investigate the region corresponding to the particular conditions of plasma: high density and high temperature. Under these conditions, (electron-ion) collisions will be, throughout this work, assumed binary and the dynamics of the electrons will be treated relativistically. Furthermore, in this work we shall use the statistical classical mechanics (not the quantum statistical mechanics as in the Fermi-Dirac distribution) because

the condition $\lambda_{th} = h/\sqrt{2\pi m_e K_B T} < N_e^{-1/3}$ (m_e is the electron mass, λ_{th} is the De Broglie thermal length and N_e is the electrons density) is fulfilled in the stated density and temperatures ranges. For example, if $N_e = 10^{24}$ cm⁻³ and $T > 10^7$ K, it is easy to verify that this inequality is correct. This condition means that the wave function extent (λ_{th}) associated with the electron is smaller than the mean distance ($\sim N_e^{-1/3}$) between two free electrons. In the present work, we focus on electron broadening in the impact approximation [1,2]. We thus, reformulate the standard semiclassical collision operator by taking into account the relativistic effects of the Lienard-Wiechert retarded electric field [3] due to the free electron movement. In addition, we assume that the plasma is optically thin (the opacity phenomenon is neglected), for this raison the spectral line shape is not influenced by the absorption process. Furthermore, by neglecting the electrons recombinaison, we assume that the decoupling of the free electrons from the radiation field is satisfied. The units system used here (unless specified) is the CGS system. In many cases in line broadening, fast particles (typically electrons) are described by a collisional approach, while particles of which field have a weak variation during the inverse half width half maximum (HWHM) time scale are considered static and treated via a quasistatic microfield. For many applications, isolated lines have great importance. So, the calculations of the broadening of such a line in a plasma are normally made by using the impact approximation for electrons [1] in the semi-classical version [4], as the ionic contribution is typically negligible.

2. Theoretical Basis of the Electron Broadening

The Stark effect is important in plasmas of high degree of ionization and high temperatures. In all cases presented in the remainder of our work, the Stark effect is dominant. The two most popular approximations in the computations of the electronic collision operator Φ are the dipole approximation and the approximation of the classical path which considers the perturbing electrons in the impact approximation. Our departure point is the expression of the intensity of the spectral line shape [2,5]:

$$I(\omega) = -\frac{1}{\pi} \operatorname{Re} \left\{ d_{\alpha\beta} \langle \langle \alpha\beta \left| \left(i\omega - \frac{i\left(H_g - H_e\right)}{\hbar} + \Phi \right)^{-1} \right| \alpha'\beta' \rangle \rangle d^*_{\alpha'\beta'} \right\}$$
(1)

where Φ is the relativistic collision operator (in Hz) which is independent from time and electric micro-field and has matrix elements given by

$$\langle \langle \alpha \beta | \Phi | \alpha' \beta' \rangle \rangle = \sum_{\alpha''} r_{\alpha \alpha''} r_{\alpha'' \alpha'} \Phi_d \left(\omega_{\alpha \alpha''}, \omega_{\alpha'' \alpha'} \right)$$

+
$$\sum_{\beta''} r_{\beta \beta''} r_{\beta'' \beta'} \Phi_d \left(\omega_{\beta \beta''}, \omega_{\beta'' \beta'} \right) - r_{\alpha \alpha'} r_{\beta' \beta} \Phi_{int} \left(\omega_{\alpha \alpha'}, \omega_{\beta' \beta} \right)$$
(2)

where α , β are the upper and lower levels respectively and r_{ab} (d_{ab} is dipole operator) is the matrix element of position operator of the bounded electron. We now aim to calculate the direct relativistic term Φ_d (in Hz/cm²) and the relativistic term of interference $\Phi_{int} = 2\Phi_d$ [5]. Before starting this task, let us remind that the criterion of validity of the impact theory, according to Voslamber [6], is not applicable for any pair (ω_1 , ω_2). However, for the isolated lines ($\omega_1 = -\omega_2$), this theoretical problem does not arise, and impact theory is valid. Specifically, the study of Φ_d is performed in the case of isolated lines for an ionic emitter and hyperbolic paths for free electrons. This treatment is based on the results obtained under the same conditions in the non-relativistic case. The relativistic collision operator is then given by

$$\Phi_d(\omega_1,\omega_2) = -\frac{2\pi N_c e^2}{3\hbar^2} \int_0^c v f(\beta) d\beta \int_{\rho_{\min}}^{\rho_{\max}} \rho_0 d\rho_0 \int_{-\infty}^{+\infty} dt_1 \int_{-\infty}^{t_1} dt_2 e^{(i\omega_1 t_1 + i\omega_2 t_2)} \left[E_{LW}(t_1) \cdot E_{LW}(t_2) \right]$$
(3)

where c is the speed of light in vacuum, v is the initial velocity of the colliding electron and $f(\beta) d\beta$ is the distribution of the velocities of Juttner-Maxwell (generalized Maxwell distribution of the velocities when the movement of particles is relativistic) given by

$$f(\beta) d\beta = \frac{\gamma^5 \beta^2 d\beta}{\theta K_2 (1/\theta)} \exp\left(-\gamma/\theta\right) \tag{4}$$

where

$$\theta = \frac{K_B T}{m_e c^2}, \quad \gamma = 1/\sqrt{(1-\beta^2)}, \quad \beta = v/c \tag{5}$$

and $K_2(1/\theta)$ is a Bessel function, *T* is the temperature, m_e is the electron mass and ρ_0 is the impact parameter, whereas ρ_{\min} and ρ_{\max} are the limits of the last integral (Formula (3)) that will be chosen later. The electric field of Lienard-Wiechert is given by [3]

$$\boldsymbol{E}_{LW}\left(\boldsymbol{R},\boldsymbol{\alpha},t\right) = -e\frac{\left(\boldsymbol{\eta}-\boldsymbol{\alpha}\right)\left(1-\boldsymbol{\alpha}^{2}\right)}{k^{3}R^{2}(t')} - \frac{e}{c^{2}}\frac{\boldsymbol{\eta}}{k^{3}R(t')} \times \left\{\left(\boldsymbol{\eta}-\boldsymbol{\alpha}\right)\times\frac{d\boldsymbol{\upsilon}(t')}{dt'}\right\}$$
(6)

then

$$\boldsymbol{\alpha} = \frac{\boldsymbol{v}\left(t'\right)}{c}, \boldsymbol{\eta} = \frac{\boldsymbol{R}(t')}{\boldsymbol{R}(t')}, t' = t - \boldsymbol{R}(t')/c \tag{7}$$

where the retarded time is given by: $t' = t - \frac{R(t')}{c}$, *e* is the charge of the electron, R(t') is the electron position vector, and $\eta = \frac{R(t')}{R(t')}$ is a unit vector directed from the position of the moving charge (electron) towards the observation point (where the emitter is located), and *k* is given by

$$k = \frac{dt}{dt'} = 1 + \frac{1}{c} \frac{dR(t')}{dt'} = 1 + \eta \alpha$$
(8)

The first term of the field (6), the velocity field, goes to the known Coulomb field when $v \rightarrow 0$ whereas the second term of the field, is the acceleration field or the radiation field. As the ratio of second term (the radiation field) of the field E_{LW} on the first term is less than v^2/c^2 , we can therefore neglect the second and use only the first part of the field in the subsequent development

$$E_{LW}(\mathbf{R},t) \simeq e\left[\frac{(\eta - \alpha)\left(1 - \alpha^{2}\right)}{k^{3}R^{2}}\right]$$
(9)

By using the approximation $1 - \alpha^2 \simeq 1$, which is justified in our subsequent studies ($T = 8 \times 10^8$ K, the probable α is about 0.22), therefore the electric field becomes

$$E_{LW}(\mathbf{R},t) = e\left[\frac{(\eta - \alpha)}{k^3 R^2}\right]$$
(10)

If we neglect the fine structure ($\omega_1 = \omega_2 = 0$), we can write the collision operator as

$$\Phi_d(0,0) = -\frac{\pi N_e e^2}{3\hbar^2} \int_0^c v f(\beta) d\beta \int_{\rho_{\min}}^{\rho_{\max}} \rho_0 d\rho_0 \int_{-\infty}^{+\infty} E_{LW}(t_1) dt_1 \int_{-\infty}^{+\infty} E_{LW}(t_2) dt_2$$
(11)

or equivalently

$$\Phi_d(0,0) = -\frac{\pi N_e e^2}{3\hbar^2} \int_0^c v f(\beta) d\beta \int_{\rho_{\min}}^{\rho_{\max}} \rho_0 d\rho_0 G^2$$
(12)

such as

$$G = -e \int_{-\infty}^{+\infty} \frac{\left(\frac{R(t')}{R(t')} - \frac{v(t')}{c}\right)}{\left(\frac{dt}{dt'}\right)k^2R^2(t')} dt$$
(13)

or, by integrating over t'

$$G = -e \int_{-\infty}^{+\infty} \frac{\left(\frac{R(t')}{R(t')} - \frac{dR(t')}{cdt'}\right)}{\left(1 + \frac{1}{c}\frac{dR(t')}{dt'}\right)^2 R^2(t')} dt'$$
(14)

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$$G = -e \int_{-\infty}^{+\infty} \left(\frac{\mathbf{R}(t')}{\left(1 + \frac{1}{c} \frac{dR(t')}{dt'}\right)^2 R^3(t')} - \frac{d\mathbf{R}(t')}{\left(1 + \frac{1}{c} \frac{dR(t')}{dt'}\right)^2 cdt' R^2(t')} \right) dt'$$
(15)

In the following, we use the notations and the variable change

$$\begin{split} \varepsilon &= (1 + \frac{m^2 v^4 \rho_0^2}{Z_{em}^2 e^4})^{1/2} \\ t' &= \frac{\rho_0}{v} \left(\epsilon \sinh\left(x\right) - x\right) \\ dt' &= \frac{\rho_0}{v} \left(\epsilon \cosh\left(x\right) - 1\right) dx \\ R\left(t'\right) &= \rho_0 \left(\epsilon \cosh\left(x\right) - 1\right) dx \\ R\left(t'\right) &= \rho_0 \left(\epsilon - \cosh\left(x\right)\right) \\ \frac{dX}{dt'} &= \frac{\frac{dX}{dx}}{\frac{dt'}{dx}} = -v \frac{\sinh(x)}{\left(\epsilon \cosh\left(x\right) - 1\right)} \\ Y &= \rho_0 \sqrt{\epsilon^2 - 1} \sinh\left(x\right) \\ \frac{dY}{dt'} &= \frac{\frac{dY}{dx}}{\frac{dt'}{dx}} = v \frac{\sqrt{\epsilon^2 - 1} \cosh\left(x\right)}{\left(\epsilon \cosh\left(x\right) - 1\right)} \\ R\left(t'\right) &= Xi + Yj \\ v\left(t'\right) &= \frac{dX}{dt'} i + \frac{dY}{dt'}j \end{split}$$

where Z_{em} is the charge of the ionic emitter and (i, j) are the basis of the cartesian coordinates. Then the function *G* is given by

$$G = -\frac{e}{\rho_0^2} \int_{-\infty}^{+\infty} \frac{X(t')\mathbf{i} + Y(t')\mathbf{j}}{\left[\left(\epsilon\cosh\left(x\right) - 1 + \frac{v}{c}\epsilon\sinh\left(x\right)\right)\right]^2 R\left(t'\right)} dt' -\frac{e}{c\rho_0^2} \int_{-\infty}^{+\infty} \frac{\frac{dX}{dt'}\mathbf{i} + \frac{dY}{dt'}\mathbf{j}}{\left[\left(\epsilon\cosh\left(x\right) - 1 + \frac{v}{c}\epsilon\sinh\left(x\right)\right)\right]^2} dt'$$
(16)

Using (17), we jump to integrate over x:

$$G = -\frac{e}{v\rho_0^2} \int_{-\infty}^{+\infty} \frac{\rho_0 \left(\epsilon - \cosh\left(x\right)\right)i + \rho_0 \sqrt{\epsilon^2 - 1} \sinh\left(x\right)j}{\left[\epsilon \cosh\left(x\right) - 1 + \frac{v}{c}\epsilon \sinh\left(x\right)\right]^2} dx$$

$$-\frac{e}{c\rho_0^2} \int_{-\infty}^{+\infty} \frac{-v \frac{\sinh(x)}{(\epsilon \cosh\left(x\right) - 1)}i + v \frac{\sqrt{\epsilon^2 - 1} \cosh\left(x\right)}{(\epsilon \cosh\left(x\right) - 1)}j}{\left[\epsilon \cosh\left(x\right) - 1 + \frac{v}{c}\epsilon \sinh\left(x\right)\right]^2} \frac{\rho_0}{v} \left(\epsilon \cosh\left(x\right) - 1\right) dx$$

$$= -\frac{e}{v\rho_0} \int_{-\infty}^{+\infty} \frac{\left(\epsilon - \cosh\left(x\right)\right)i + \sqrt{\epsilon^2 - 1} \sinh\left(x\right)j}{\left[\epsilon \cosh\left(x\right) - 1 + \frac{v}{c}\epsilon \sinh\left(x\right)\right]^2} dx$$

$$-\frac{e}{c\rho_0} \int_{-\infty}^{+\infty} \frac{-\sinh(x)i + \sqrt{\epsilon^2 - 1} \cosh\left(x\right)j}{\left[\epsilon \cosh\left(x\right) - 1 + \frac{v}{c}\epsilon \sinh\left(x\right)\right]^2} dx$$
(17)

or in a more simplified form

$$\Phi_d(0,0) = -\frac{N_e e^4}{3\pi\hbar^2} \int_0^c \frac{\gamma^5 \beta d\beta}{\theta c K_2 (1/\theta)} \exp\left(-\gamma/\theta\right) \int_{\rho_{\min}}^{\rho_{\max}} \frac{d\rho_0}{\rho_0} \left[\overline{A}^2 + (\epsilon^2 - 1)\overline{B}^2\right]$$
(18)

where

$$\overline{A} = \int_{-\infty}^{+\infty} \frac{-(\epsilon - \cosh(x)) + \beta \sinh(x)}{\left[\epsilon \cosh(x) - 1 + \beta\epsilon \sinh(x)\right]^2} dx$$
$$= 2 \frac{(\epsilon^2 - 1)\sqrt{1 - \epsilon^2 + \beta^2 \epsilon^2} - \beta^2 \epsilon^2 \tanh^{-1}(\sqrt{1 - \epsilon^2 + \beta^2 \epsilon^2})}{\epsilon (1 - \epsilon^2 + \beta^2 \epsilon^2)^{3/2}}$$
(19)

$$\overline{B} = -\int_{-\infty}^{+\infty} \frac{\sinh(x) + \beta \cosh(x)}{\left[\epsilon \cosh(x) - 1 + \beta \epsilon \sinh(x)\right]^2} dx = 0$$
⁽²⁰⁾

Then the relativistic collision operator caused by the Lienard-Wiechert electric field $\Phi_{d,LW}$ is given by

$$\Phi_{d}(0,0)(Hz/cm^{2}) = \Phi_{d,LW} = -\frac{4N_{e}e^{4}}{3\pi\hbar^{2}\theta cK_{2}(1/\theta)} \int_{0}^{1} \frac{\exp(-\frac{1}{\theta\sqrt{1-\beta^{2}}})}{(1-\beta^{2})^{5/2}} \beta d\beta \int_{\rho_{\min}}^{\rho_{\max}} \frac{[(\epsilon^{2}-1)\delta-\beta^{2}\epsilon^{2}\tanh^{-1}(\delta)]^{2}}{\epsilon^{2}\rho_{0}\delta^{6}} d\rho_{0}$$

$$(21)$$

where we have put

$$\delta \sim \delta(\beta, \epsilon) = \sqrt{1 - \epsilon^2 + \beta^2 \epsilon^2}$$
(22)

By taking the maximum of the impact parameter $\rho_{max} = 0.68\lambda_D$ (λ_D is the Debye length) [5] and the minimum of the impact parameter equal to Bohr radius (the Wiesskopf radius is much smaller than Bohr radius in our application), and after numerical integration of (31) we find the following result in Table 1.

We note that we have considered the lower limit of the integration over the impact parameter equal to the Bohr radius, because in our study we only intented to compare the two collision operators corresponding to Coulomb and Lienard-Wiechert interactions. In reality, as Formula (31) shows, by decreasing the lower limit of the impact parameter, the value of the collision operator increases and by increasing the lower limit, the value of the collision operator decreases. Another reason to mention is: by regarding Formula (3) of Ref. [5], we see that the minimum of the impact parameter (Wiesskopf radius r_W), in our conditions of the high temperature, high charge number Z = 70 and the upper and lower levels of the transition $n_a = 3$ (1s3d), $n_b = 2$ (1s2p)(in triplet case), is much smaller than the Bohr radius a_0 that is to say $r_W << a_0$. For example, for a density 10^{18} cm⁻³, for the lower limit r_W , the value of the collision operator $\Phi_{d,LW}$ is 0.35×10^{-2} eV (that is overestimated in our opinion) whereas it is 0.21×10^{-2} eV for the lower limit equal to the Bohr radius a_0 . In fact, in the region $r < a_0$, the quantum effects must be taken into account. For these reasons, we have only considered, in our calculation, the minimum of the impact parameter equal to the Bohr radius a_0 . In addition, we remark from the Table 1 that the effect of the Lienard-Wiechert field is to reduce the amplitude of the collision operator. This reduction is more pronouced for the weak electron densities. We must also note that the criteria of the isolated lines becomes not valid for densities great than 10^{21} cm⁻³ because the FWHM = $2 \times$ HWHM of two neighboring spectral lines becomes of the same order of magnitude of the separation (about 3eV) between the line arising from 1s3d to 1s2p (triplet case that we study) transition and the neighboring line arising from 1s3p to 1s2s (singlet case) transition. Following the Table 1, and for densities less than 10^{21} cm⁻³, the HWHM is small enough to be can considered that the studied line is isolated.

Table 1. Comparison between collision operators for Coulomb interaction $\Phi_{d,C}$ and for Lienard Wiechert interaction $\Phi_{d,LW}$ (multiplied by \hbar in eV*s and by a_0^2 in cm² where a_0 is Bohr radius) at a temperature $T = 8 \times 10^8$ K and different densities and for Helium-like Ytterbium $Z_{em} = 68$, for the radiative transition 1s3d to 1s2p for triplet case. After this multiplication, the following results are in eV.

Ne in $\rm cm^{-3}$	$\Phi_{d,C}$ in eV	$\Phi_{d,LW}$ in eV	$HWHM_C$ in eV	HWHM_{LW} in eV	Percent
10^{16}	$0.32 imes 10^{-4}$	$0.24 imes 10^{-4}$	$0.25 imes 10^{-4}$	0.17×10^{-4}	25
10^{18}	$0.27 imes 10^{-2}$	$0.21 imes 10^{-2}$	$0.20 imes 10^{-2}$	$0.15 imes10^{-2}$	22
10^{20}	0.24	0.22	0.18	0.17	8
10 ²²	26	25	20	19	4

Note: We have defined the percent to be equal $\left(\frac{\Phi_{d,C}-\Phi_{d,LW}}{\Phi_{d,C}}\cdot 100\right)$.

3. Conclusions

In this work, we have investigated Lienard-Wiechert or retarded electric fields produced by moving electric charges with respect to a rest frame. Specifically, we have studied its contribution to the broadening of the spectral line shape of the Helium-like Ytterbium in hot and dense plasmas radiative (transition from 1s3d to 1s2p). The principal result, as the table shows is: the retarded Lienard-Wiechert interaction, narrows the line shape comparatively to the pure Coulomb interaction. The narrowing is more pronouced for the low densities.

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Article



Using the Pairs of Lines Broadened by Collisions with Neutral and Charged Particles for Gas Temperature Determination of Argon Non-Thermal Plasmas at Atmospheric Pressure

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Abstract: The spectroscopic method for gas temperature determination in argon non-thermal plasmas sustained at atmospheric pressure proposed recently by *Spectrochimica Acta Part B* 129 14 (2017)—based on collisional broadening measurements of selected pairs of argon atomic lines, has been applied to other pairs of argon atomic lines, and the discrepancies found in some of these results have been analyzed. For validation purposes, the values of the gas temperature obtained using the different pairs of lines have been compared with the rotational temperatures derived from the OH ro-vibrational bands, using the Boltzmann-plot technique.

Keywords: plasma spectroscopy; microwave discharges; gas temperature; stark broadening parameters; atomic emission spectroscopy

1. Introduction

Optical emission spectroscopy (OES) techniques based on the analysis of molecular emission spectra are commonly used for gas temperature (T_g) determination of plasmas at atmospheric pressure. The rotational temperature derived from them is considered as a good estimation of the kinetic temperature of the plasma heavy particles [1,2] thanks to the strong coupling between translational and rotational energy states under high-pressure conditions. The emissions of diatomic species, such as OH, N₂, N₂⁺, CN, ..., have been traditionally employed with this purpose [3–10], but the use of molecular emission spectroscopy is not always easy for gas temperature measurement in plasmas: overlapping of bands, rotational population distribution of levels having a non-Boltzmann nature, wake emission of rotational bands, among others, can make it difficult to obtain reliable values of gas temperature.

Alternative OES methods for gas temperature determination are needed. In this way, the van der Waals broadening of some atomic lines has been used for this purpose due to its dependence on the plasma gas temperature. For argon plasmas, the 425.9, 522.1, 549.6, and 603.2 nm argon lines have been the most frequently employed [11–14]. This technique is based on the detection of argon lines not affected by resonance broadening (also related to T_g). It requires the use of additional techniques for simultaneous determination of the electron density [15], as these lines have a non-negligible Stark broadening for electron densities above 10^{14} cm⁻³, which needs to be determined. Yubero et al. in [16] proposed a method to circumvent this dependence on electron density by considering pairs of these

lines. This method allows for the determination of the gas temperature from the measurements of Lorentzian profiles of some pairs of argon atomic lines, and when applying it, no assumptions on the degree of thermodynamic equilibrium among excited states are needed (unlike methods based on rotational temperature determination).

However, the authors of [16] found a small disagreement in the results from pairs Ar I 603.2 nm/Ar I 522.1nm and Ar I 549.6 nm/Ar I 522.1 nm. In the present work, this disagreement is explained, and other pairs of lines have been employed for the gas temperature determination.

2. Method

This method is valid for plasmas sustained at atmospheric pressure whose line profiles can be fitted to a Voigt function-characterized by a full-width at half-maximum (FWHM), W_V -resulting from the convolution of a Gaussian function (W_G) with a Lorentzian function (W_L) (see, e.g., [17,18]). Indeed, the profiles of atomic lines emitted by plasmas with no presence of magnetic fields result from different broadening mechanisms leading to Gaussian or Lorentzian profile shapes, briefly described below.

The motion of emitting atoms with respect to the detector, with a continuous velocity distribution depending on their temperature, leads to the so called *Doppler broadening* and a Gaussian-shaped line profile with a FMHW W_D (in nm) given by

$$W_D = 7.16 \cdot 10^{-7} \lambda \sqrt{T_g/M} \tag{1}$$

where λ , T_g , and M are the wavelength (nm), the gas temperature (in K), and the mass of the radiating atom (in a.m.u.), respectively.

The *van der Waals broadening* is due to the dipole moment induced by neutral perturber atoms interacting with the electric field of the excited emitter atom and generates line profiles with a Lorentzian shape (with an FWHM W_W), according to the Lindholm–Foley theory [19].

The *resonance broadening* of spectral lines is due to dipole–dipole interactions of the emitter with ground-state atoms of the same element [20] and contributes to the Lorentzian part of the profile with an FWHM W_R .

The *Stark broadening* (FWHM W_S) of a line is due to interactions of the emitter atom with the surrounding charged particles, perturbing the electric field it experiences. In the case of a non-hydrogenic atom, the profiles of isolated spectral lines broadened by collisions with electrons have a Lorentzian shape. For thermal plasmas with a gas temperature similar to the electron one, the mobility of ions is high and the impact approximation [21] is also valid for ions, being their contribution to the broadening also being Lorentzian. In the ion impact limit, line profiles are symmetric Lorentzian. On the contrary, for plasmas where the ion mobility is small (e.g., plasmas with gas temperature relatively low), a quasistatic approximation is often needed to model the ion broadening in order to explain the slightly asymmetric shape of the profiles. The less dynamical the ions are, the more asymmetric the lines are. The finite lifetime of the excited levels gives rise to *natural broadening*, which is typically very small (~0.00001 nm) and can be neglected in the case of atmospheric pressure plasma spectroscopy.

Finally, the line profile is also affected by the instrumental function of the spectrometer used for its detection. Usually, this instrumental function can be well approximated by a Gaussian profile with an FWHM W_I , as shown in the next section.

Thus, the broadening contributions with a Gaussian shape will lead to a profile with an FWHM (W_G) given by

$$W_G = \sqrt{(W_D)^2 + (W_I)^2}$$
(2)

and those having a Lorentzian shape give rise to a profile that is also Lorentzian with an FWHM (W_L) given by

$$W_L = W_R + W_S + W_W. \tag{3}$$

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The method we propose here only considers atomic lines with a negligible resonance broadening. Thus, the Lorentzian part of a line profile is only due to Stark and van der Waals broadenings, and full width at half maximum of the Lorentzian profile, W_L can be written as follows:

$$W_L = W_S + W_W \tag{4}$$

Several studies of Stark broadening for atomic and singly charged ion lines [18,22–26] show that Equation (4) can be written as

$$W_S \cong \left[w_S^e + w_S^i \right] \frac{n_e}{10^{16}} \tag{5}$$

where parameters w_{S}^{e} and w_{S}^{i} are electronic and ionic contributions to the full-width at half-intensity maximum given for an electron density equal to 10^{16} cm⁻³, and n_{e} is the electron density.

In non-thermal-plasmas where T_g is much lower than T_e (electron temperature), the ionic contribution w_S^i can be neglected. Additionally, W_S can be considered to have a weak dependence on T_e in the small range of electron temperature from 5000 to 10,000 K [22]. In this way, W_S depends only on n_e and Equation (5) can be approximated as follows:

$$W_L(T_g, n_e) = w_S \frac{n_e}{10^{16}} + W_W(T_g)$$
(6)

where $w_S = w_S^e$.

On the other hand, the van der Waals broadening has a full width at half maximum W_W in nm given by Griem [18], which, considering the ideal gas equation $N = P/K_BT_g$ for the density of perturbers, and where K_B is the Boltzmann constant and P is the pressure, can be written as

$$W_W(T_g) = \frac{C}{T_g^{\frac{7}{10}}}(nm)$$
(7)

with *C* being determined by the type of gas in the discharge and the nature of the atom emitters:

$$C = \frac{8.18 \cdot 10^{-19} \lambda^2 \left(\alpha \left\langle \overline{R^2} \right\rangle \right)^{2/5} P}{k_\beta \mu^{3/10}} (\mathrm{nm} \cdot \mathrm{K}^{7/10})$$
(8)

where

$$\left\langle \overline{R^2} \right\rangle = \left\langle \overline{R_{U}^2} \right\rangle - \left\langle \overline{R_{L}^2} \right\rangle \tag{9}$$

is the difference of the squares of coordinate vectors (in a_0 units) of the upper and lower level, λ is the wavelength of the observed line in nm, α is the polarizability of perturbers interacting with the excited radiator in cm³, T_g is the temperature of the emitters (coincident with the gas temperature) in K, and μ is the reduced mass of the emitter–perturber pair in a.m.u.

For argon plasma at atmospheric pressure, when considering the van der Waals broadening of argon atomic lines ($\mu = 19.97$ and $\alpha = 16.54 \cdot 10^{-25}$ cm³), *C* can be written as [16]

$$C = 7.5 \cdot 10^{-7} \lambda^2 \left(\left\langle \overline{R^2} \right\rangle \right)^{2/5} (\operatorname{nm} \cdot K^{7/10}).$$
⁽¹⁰⁾

Thus, the formula for W_L given by Equation (3) can be approximately expressed as

$$W_L(T_g, n_e) = w_S \frac{n_e}{10^{16}} + \frac{C}{T_g^{\frac{7}{10}}}.$$
(11)

The method for gas temperature determination proposed by Yubero et al. in [16] is based on the measurement of the Lorentzian contribution to the entire FWHM for two atomic lines, *L*1 and *L*2:

$$W_L^{L1}(T_g, n_e) = w_S^{L1} \frac{n_e}{10^{16}} + \frac{C_L^{L1}}{T_g^{\frac{1}{10}}} W_L^{L2}(T_g, n_e) = w_S^{L2} \frac{n_e}{10^{16}} + \frac{C_L^{L2}}{T_g^{\frac{1}{10}}} .$$
(12)

The dependence on the electron density can be eliminated from these expressions, and a linear dependence between the FWHM of the Lorentzian part of the total profile of these two lines, W_L^{L1} and W_l^{L2} , is obtained:

$$W_L^{L1} = \frac{w_S^{L1}}{w_S^{L2}} W_L^{L2} + \left(C^{L1} - C^{L2} \frac{w_S^{L1}}{w_S^{L2}} \right) T_g^{-0.7}.$$
 (13)

Finally, the gas temperature is given by

$$T_g = \left(\frac{C^{L1} - C^{L2} \frac{w_5^{L1}}{w_5^{L2}}}{W_L^{L1} - \frac{w_5^{L1}}{w_5^{L2}}}W_L^{L2}\right)^{1/0.7}.$$
(14)

Thus, as long as the Stark parameters for two lines and their *C* parameters are known, the gas temperature can be determined from full Lorentzian FWHMs of these lines. Nevertheless, the applicability of this method relies on the knowledge of these parameters, with a certain degree of accuracy. The theoretical estimation of these parameters in the literature can lead to significant errors in gas temperature values.

Concerning this matter, Yubero et al. [16] gather the values of FWHM of the Lorentzian profile of the lines Ar I 603.2 nm, Ar I 549.6 nm, and Ar I 522.1 nm These lines correspond to transitions from high energy levels and exhibit slightly asymmetric profiles [13]. The effect of this asymmetry, not considered in the previous work [16], has been removed from the profiles according to the procedure in [13] (so, only considering the symmetric electron contribution) in order to improve the results of the method. According to this reference, the effect of ions is more important in the right shape of profile. Therefore, only the left part of the profile has been considered to generate the entire profile. Figure 1 shows an example of the symmetrization procedure for the Ar I 522.1 nm line.



Figure 1. Symmetrization of Ar I 522.1 nm.

In addition, new Ar I lines at 560.7 nm and 518.8 nm have been included in this work.

Measured values of W_L for these lines allow us to obtain T_g using Equation (14), provided the *C* coefficients and Stark broadening parameters are known. Table 1 includes values of the *C* coefficients calculated for these lines from Equation (8), and the Stark broadening parameters theoretically determined by Dimitrijević et al. [22] for an electron temperature of 10,000 K.

Table 1. *C* coefficients calculated from Equation (8), and the Stark broadening parameters due to electron impacts theoretically determined by Dimitrijević et al. [22] for an electron temperature of 10,000 K and an electron density of 10^{16} cm⁻³.

Ar I Line (nm)	С	$w_S=w^e_S$ (nm)
603.2	4.2	0.149
549.6	4.9	0.305
522.1	5.9	0.588
560.7	3.6	0.145
518.8	4.1	0.104

3. Experimental Set-Up

In this work, we measured the gas temperature of an argon microwave (2.45 GHz) induced plasma column sustained at atmospheric pressure and generated inside a quartz tube (with the inner and outer diameters 1.5 mm and 4 mm, respectively), described elsewhere [27]. A similar plasma had been previously characterized in [28], its electron density being of the order of 10^{14} (cm⁻³). The electron temperature was estimated to be close to 10,000 K from observed relative populations of the argon excited levels assuming a partial local thermodynamic equilibrium [28].

Figure 2 includes a scheme of the optical detection assembly and data acquisition system to process spectroscopic measurements. A *surfaguide* was employed as a coupling device, injecting a microwave power of 100 W to the plasma. The argon flow rate was set at 0.5 slm (standard liters per minute) and adjusted with a calibrated mass flow controller.



Figure 2. Scheme of experimental set-up.

Light emission from the plasma was analyzed by using a Czerny–Turner type spectrometer with a 1 m focal length, equipped with a 2400 grooves/mm holographic grating and a photomultiplier (spectral output interval of 200–800 mm) as a detector. The light emitted by the plasma was collected side-on using an optical fiber at different axial positions along the plasma column (z = 4, 8, and 12 cm measured from the end of the column).

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The instrumental function of the spectrometer was measured from the FWHM of the line Ne I 632.8 nm emitted by a helium–neon laser (this line is a good choice to make this estimate, as it has a wavelength close to those of the Ar I lines considered in this work). When using equal entrance and exit spectrometer slit widths, this function had an approximately triangular shape, which could be well-fitted with a Gaussian function. In this way, an instrumental broadening width $W_I = (0.032 \pm 0.001)$ nm was measured when using slit widths of 100 µm.

On the other hand, measurements of the light absorption have shown that the plasma studied can be considered as optically thin in the direction of observation chosen (transversally) for the Ar I lines detected [28,29].

Each Ar I line was measured experimentally several times and was fitted to a Voigt profile. Therefore, the uncertainty of each Voigt FWHM corresponds to the dispersion. The Lorentzian contribution to the entire broadening in each case was obtained from the Voigt FWHM measured for each line using the formula [30,31]

$$W_V \approx \frac{W_L}{2} + \sqrt{\left(\frac{W_L}{2}\right)^2 + W_G^2} \tag{15}$$

assuming that $W_G \approx W_I$, since, according to Equation (2), the Doppler contribution can be considered as negligible when compared to the instrumental one under the experimental conditions in the plasma studied ($T_g \leq 2500$ K, W_D ^{ArI} ≤ 0.003 nm).

4. Results

Table 2 shows the values of Lorentzian FWHM measured for the different Ar I lines at different positions. For each pair of lines, these values allow for T_g determination using Equation (14), provided *C* coefficients and Stark broadening parameters given in Table 1.

Table 2. Lorentzian FWHM of lines Ar I 603.2 nm, Ar I 549.6 nm, Ar I 522.1 nm, Ar I 560.7 nm, and Ar I 518.8 nm measured at different axial plasma positions.

<i>z</i> (cm)	z = 4 cm	z = 8 cm	<i>z</i> = 12 cm
W_{L}^{603} (nm)	0.0411 ± 0.0014	0.0437 ± 0.0016	0.0459 ± 0.0014
W_L^{549} (nm)	0.0594 ± 0.0019	0.0626 ± 0.0016	0.0731 ± 0.0012
W_L^{522} (nm)	0.0958 ± 0.0024	0.1020 ± 0.0019	0.122 ± 0.002
W_L^{560} (nm)	0.0342 ± 0.0024	0.0377 ± 0.0024	0.0429 ± 0.0012
W_L^{518} (nm)	0.0372 ± 0.0021	0.0418 ± 0.0016	0.0472 ± 0.0018

Table 3 shows the T_g values obtained using these parameters for an electron temperature of 10,000 K. These values are also compared with the values obtained using OH ro-vibrational band [13]. Uncertainties in T_g have been obtained from Equation (14) by considering *C* and W_S as theoretical constants and only taking into account uncertainties in the broadenings experimentally measured and errors of approximations used in this method.

Overall, results from the method of pairs of argon lines proposed in this work are in good agreement with those derived from the analysis of OH ro-vibrational band. However, there are some results affected by a very high uncertainty (higher than 50%) corresponding, in some cases, to the Ar I 560.7 nm line. Additionally, there are values lower than the ones obtained from rotational temperature technique corresponding mainly to the 518.8 line.

Table 3. Gas temperature obtained using Equation (14) with theoretical Stark broadening parameters
at electron temperatures of 10,000 K, and comparison with the one obtained from OH ro-vibrational
bands, using the well known Boltzmann plot technique.

$z = 4 \text{ cm } T^{BP}_{g}$ (K) = 1390 ± 70	Ar I 549 nm	Ar I 522 nm	Ar I 560 nm	Ar I 518 nm
Ar I 603 nm	1300 ± 300	1420 ± 180	570 ± 240	1100 ± 500
Ar I 549 nm	-	1800 ± 900	2200 ± 1300	1200 ± 230
Ar I 522 nm		-	2000 ± 700	1300 ± 200
Ar I 560 nm			-	900 ± 300
$z = 8 \text{ cm } T^{BP}_{g}$ (K) = 1330 ± 70	Ar I 549 nm	Ar I 522 nm	Ar I 560 nm	Ar I 518 nm
Ar I 603 nm	1100 ± 220	1300 ± 200	700 ± 600	760 ± 220
Ar I 549 nm	-	1700 ± 500	1400 ± 500	920 ± 120
Ar I 522 nm		-	1600 ± 500	1030 ± 110
Ar I 560 nm			-	800 ± 300
$z = 12 \text{ cm } T^{BP}{}_g$ (K) = 1520 \pm 70	Ar I 549 nm	Ar I 522 nm	Ar I 560 nm	Ar I 518 nm
Ar I 603 nm	1600 ± 400	1680 ± 220	3000 ± 3000	500 ± 130
Ar I 549 nm	-	1800 ± 500	1400 ± 400	810 ± 110
Ar I 522 nm		-	1500 ± 300	930 ± 110
Ar I 560 nm			-	640 ± 130

5. Discussion and Conclusions

In this paper, the OES tool proposed by Yubero et al. in [16] for the determination of the gas temperature in non-thermal plasmas, based on the measurement of the Lorentzian part of the profile of a pair of atomic emission lines, has been improved, as asymmetries of the profiles of these lines have been removed. Compared with the results of [16], better agreement with gas temperatures obtained from OH ro-vibrotional band has been found. We can conclude that symmetrization of the line profile is advisable to obtain good results from this method. Moreover, results from new lines Ar I 560.7 nm and Ar I 518.8 nm have been also included in this paper. As can be seen in Table 3, there are some pairs that do not give accurate results or T_g values with higher uncertainty. They correspond to those cases in which the denominator in Equation (14) is very small, so gas temperature determination becomes very sensitive, giving rise to large errors. Examples of pairs of line giving large errors are Ar I 560.7 nm/Ar I 603.2 nm and Ar I 518.8 nm/Ar I 509.7 nm/Ar I 518.8 nm/Ar I 560.7 nm.

Although the method has been developed for some pairs of argon lines, it also applies for any pair of atomic lines, as long as they fulfill the following conditions:

(i) Their Stark parameters must be as accurate as possible; (ii) They have a negligible resonance broadening (or they do not have any at all). Choosing lines with an upper or lower level that do not have an electric dipole transition (resonance line) to the ground state is a way to ensure that this condition is satisfied. Additionally, analytical expressions for resonant FWHM in the literature [32,33] can be used to evaluate the importance of this broadening; (iii) Their Stark and van der Waals broadenings should not be very different from each other, so as to avoid large errors in T_g determination (see Equation (14)); this condition applies to plasmas with a relatively low gas temperature (van der Waals broadening not negligible) and moderate electron densities (significant Stark broadening).

The method we propose in this work can be considered a good alternative to the traditional ones based on the measurement of rotational temperatures. In the application of this method, no assumptions on the degree of thermodynamic equilibrium for excited states existing in the plasma are needed, which is its main advantage compared to other methods.

Additionally, as other optical emission spectroscopy techniques, this is a non-plasma diagnosis method, easy to implement.

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Article



Effect of Turbulence on Line Shapes in Astrophysical and Fusion Plasmas

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Abstract: We look at the effect of wave collapse turbulence on a hydrogen line shape in plasma. An atom immersed in plasma affected by strong Langmuir turbulence may be perturbed by a sequence of wave packets with a maximum electric field magnitude that is larger than the Holtsmark field. For such conditions, we propose to calculate the shape of the hydrogen Lyman α Lyman β and Balmer α lines with a numerical integration of the Schrödinger equation coupled to a simulation of a sequence of electric fields modeling the effects of the Langmuir wave. We present and discuss several line profiles of Lyman and Balmer lines.

Keywords: line shape; wave collapse; electric field solitons; plasma turbulence

1. Introduction

The problem of plasma turbulence is of interest both from a theoretical point of view and from an experimental one for laboratory, fusion, and astrophysical plasmas. Plasma turbulence affects the transport and radiation properties of many kinds of plasma. In magnetic fusion studies, the quality of the plasma confinement is strongly dependent on the level of turbulent fluctuations. The first observations of turbulent fluctuations have been made in astrophysics on line shapes dominated by the Doppler effect [1,2]. If nonthermal movements take place on the line of sight, the line shape no longer corresponds to a Maxwellian velocity distribution at the emitter temperature. In the simplest models, a nonthermal velocity is defined as one that allows a quantitative measure of turbulence. The study of line shapes may then provide valuable information on the nature of turbulence, and this has been used in astrophysical and laboratory plasmas [3]. In this work, we are mainly interested in the contribution of Stark effect to the line shapes of hydrogen plasmas. The turbulent fluctuations of the plasma are created by the instabilities appearing in the different types of plasma studied [3].

One kind of plasma turbulence suspected to be present in astrophysical and fusion plasma is driven by plasma waves and electromagnetic waves. We studied the case of nonlinear wave collapse turbulence, a phenomenon occurring in the presence of an external source of energy, and coupling nonlinearly to the Langmuir waves with ion sound and electromagnetic waves. Due to this coupling, the density fluctuations associated with ion sound waves refracts the Langmuir waves in regions of low densities. Coherent wave packets localize in such regions, and experience a cycle driven by the ponderomotive force, which decreases them to shorter scales and enhances their intensity (wave collapse). In such conditions, numerous wave collapse sites are present in the plasma, which change its radiative properties. We proposed a model for calculating the change in the line shape of atoms submitted to the electric field of a nearby wave collapse [3,4]. Our model uses the numerical
solution of the emitter Schrödinger equation submitted to an electric field taken as a sequence of envelope solitons oscillating at the plasma frequency. We used the results of numerical simulations of wave collapse [5] to sample the lifetime of each soliton as well as the probability density function for the magnitude of the electric field. We will present the changes expected on a line shape of hydrogen for plasma conditions of interest in astrophysical and fusion plasmas.

The aim of this work is to study the effect of wave collapse on spectral line shapes of Lyman α , Lyman β , and Balmer α emitted by hydrogen atoms. In the following, we consider an atom submitted to a sequence of an electric field modulated by an envelope soliton, and we calculate numerically the atomic dipole autocorrelation function in the single presence of such solitons. Plasmas submitted to an energetic beam of particles or to a strong radiation can be found in many situations. In an astrophysical context, active galactic nuclei [6], pulsar radio sources [7], planetary foreshocks [8], or solar type III radio bursts [9] are possible candidates. Relevant laboratory plasmas are laser plasmas [10], radio experiments [11], or possibly also magnetic fusion plasma, since such plasmas can be affected by the energetic beams of runaway electrons [12].

The paper is organized as follows: in Section 2, we recall the main properties of turbulent Langmuir fields, and we propose a model for computing the dipole autocorrelation function (DAF) and the line shape in Section 3. We present and discuss our results on the hydrogen lines in Section 4.

2. Wave Collapse and Strong Turbulence

The physics of Langmuir turbulence have been studied in detail, since it is necessary to the understanding many radiative and transport properties of plasma. Langmuir turbulence describes a plasma state affected by a high level of excited Langmuir waves [13]. Using the Zakharov equations [14], it is possible to distinguish between weak and strong Langmuir turbulence. A useful quantity for making such a distinction is the ratio *W* of the wave energy density to the thermal energy density:

$$W = \varepsilon_0 E_L^2 / 4N_e k_B T, \tag{1}$$

with T and N_e representing the hydrogen plasma temperature and density, E_L the magnitude of the wave, k_B the Boltzmann constant, and ε_0 the permittivity of free space. Beyond a critical value of W, which depends on the plasma conditions, the Langmuir waves couple with ion sound and electromagnetic waves, resulting in a strong turbulence regime. Strong Langmuir turbulence occurs in plasma submitted to an external source of energy, which may be coupled to the plasma waves, thus increasing their intensity, and allowing the start of nonlinear processes such as wave-wave interactions [15]. In this strong turbulence regime, the energy density of the waves can exceed the plasma energy density, and a large amount of energy is available. The physical process at work is the creation of low density regions by the coupling of the density fluctuations associated to the ion sound wave with the Langmuir wave. Wave packets are refracted in regions of low density, which are also regions of high refractive index. The nonlinear ponderomotive force then moves part of the plasma out of the region of maximum field value, thus starting a dynamic process where coherent wave packets evolve to shorter scales and higher intensities reaching several hundred times the average microfield $E_0 = 1/(4\pi\epsilon_0 r_0^2)$, where r_0 is the average distance between particles defined by $r_0^3 = 3/(4\pi N_e)$. Plasma computer simulations reveal the existence of a wave packet cycle with a collapse arrested by dissipation, and a nucleation mechanism allowing the creation of new wave packets [5]. The electric field of such wave packets oscillates at a frequency close to the plasma frequency $\omega_p = \sqrt{N_e e^2/m\varepsilon_0}$, where *m* is the electron mass, and the wave packet is modulated by an envelope soliton with a Gaussian or Lorentzian shape. The average duration of a cycle is an estimate for the characteristic time of strong turbulence. It can be obtained from plasma simulations such as particle in cells codes and scales as much as 40 times the inverse of the average of W [5], using units of the inverse plasma frequency. Taking account of this relation and of the expression of W given by Equation (1), the choice of a value of W also determines the values of the electric field modulus and of the average duration of a cycle for

given background plasma conditions. If energy is supplied from an external source, localized wave packets may be created at a high rate, and the plasma will contain many of those coexisting wave packets. The localized wave packets appear to be densely packed, so that a large number of emitters experience the field of a wave packet.

3. Line Shape Model for Wave Collapse

In this work, we are interested in studying the effect of nonlinear wave collapse on a Stark spectral line shape of hydrogen atoms. Following our description of wave packet collapse, we assume that a large number of hydrogen atoms are submitted to the electric field of a wave packet. We propose to model the electric field of the wave by a sequence of solitons using a renewal process. The maximum magnitude of the electric field is sampled with a probability density function (PDF) that we assume to be half-normal. We jump from one soliton to the next using an exponential waiting time distribution $\nu \exp(-\nu t)$, with the jumping frequency ν chosen as the inverse of the average duration of a cycle [3,4]. We call such a sequence of envelope solitons a single electric field history with a duration of the order of the line shape time of interest (inverse of the line width). This time is also the decorrelation time of dipole autocorrelation function (DAF), a quantity *C*(*t*) defined by [16]:

$$C(t) = Tr\langle \vec{D} \cdot U^+(t)\vec{D}U(t)\rho\rangle, \qquad (2)$$

where the trace is over the atomic states, D and U are the atomic dipole and evolution operators, ρ is the density matrix, and the angle brackets imply an average over the configurations of the perturbation. The atom perturbation dynamics are obtained by numerically solving the Schrödinger equation for the evolution operator U(t) for each history. The DAF is obtained as an average over a large number (10^4) of independent electric field histories. The sampling of the stochastic variables may be done on the computer with pseudorandom number algorithms, associated to numerical techniques such as transformation or rejection methods [17,18]. The line shape is also obtained numerically using a Fourier transform of the DAF:

$$I(\omega) = \frac{1}{\pi} Re \int_0^\infty \exp(i\omega t) C(t) dt.$$
 (3)

The calculation presented in the following concerns the hydrogen Lyman α (L $_{\alpha}$), Lyman β (L $_{\beta}$), and Balmer α (H $_{\alpha}$) lines, neglecting fine structure in order to obtain a fast numerical evaluation, and using the spherical quantum number n, l, m. For our calculations of the Balmer lines, we neglect the broadening of the lower states of the transitions. Using our simulation, it is possible to calculate the effect of Langmuir solitons alone. It is possible to compare to Stark broadening (impact approximation or ab initio) in a hydrogen plasma in equilibrium [16]. Using a convolution, it is also possible to calculate a profile taking into account both equilibrium Stark broadening and Langmuir solitons.

4. Results

We first compute the solution of the Schrödinger equation in the presence of the sequence of solitons alone. We calculate the dipole autocorrelation function C(t) and the line profile for L_{α} , L_{β} , and H_{α} , for a density of 10^{19} m^{-3} and for a temperature of 10^5 K , conditions which can be found in the edge of a tokamak plasma. Solitons are generated for an average value of $W \approx 1.1$, resulting in a jumping frequency of $\nu \approx \omega_p/37$ and an average peak value of $150E_0$ for the electric field. After a study of the shape effect of the shape of the envelope soliton [3], and with the help of plasma simulations, we chose a soliton shape with a width equal to 20% of the wave cycle duration, a value which minimizes the broadening effect of the soliton sequence [3].

Figures 1–3 plot the DAF of L_{α} , L_{β} , and H_{α} for the pure Stark effect using an impact approximation (dashed line), pure soliton effect (solid line), and the product of the two preceding DAFs (dotted line). In Figure 1, a strong decay is observed on the soliton DAF of L_{α} (solid line)

for times shorter than the average duration of a wave packet cycle. This strong initial decay can be attributed to the large magnitude of the soliton electric field. A similar behavior of the DAF for short times is also observed for L_{β} (Figure 2, solid line) and H_{α} (Figure 3, solid line). For intermediate times, the soliton DAF of L_{α} has a weaker decay than the Stark DAF, but the two curves are similar for long times. For times longer than the wave packet cycle, we observe for L_{β} and H_{α} a weaker decay of the soliton than the impact DAF. Small amplitude oscillations are seen on these DAF, and are in phase with the plasma frequency. In all cases, the decay of the product DAF (dotted line) is significantly larger than for the Stark DAF.



Figure 1. Dipole autocorrelation functions (DAF) for L_{α} submitted to 10⁴ solitons sequences calculated for W = 1.1 (solid line) for 10,000 histories, and compared to the Stark DAF (dashed line) calculated with an impact approximation, and the product DAF (dotted line) in plasma with a density of $N_e = 10^{19} \text{ m}^{-3}$ and temperature of $T_e = 10^5 \text{ K}$.



Figure 2. Same as Figure 1 for L_{β} .

In Figures 4–6, we present the line shapes of L_{α} , L_{β} , and H_{α} for the same plasma conditions. Although being especially studied for high density plasmas, detailed Stark line shapes are also needed for low density plasmas, since they enter in the modeling of radiative transfer together with Doppler broadening [19]. We calculated the line shape profile of L_{α} , L_{β} , and H_{α} by a Fourier transform of the dipole autocorrelation functions already discussed. In Figure 4, we found that the line profile which is only affected by solitons (solid line) is similar to the Stark profile (dashed line) for L_{α} . The convolution profile (dotted line) is, however, about 2.1 broader than the Stark profile. For L_{β} (Figure 5) and H_{α} (Figure 6), the profile only affected by solitons (solid line) is narrower than the pure Stark profile. The convolution profile (dotted line) is broader by a factor 2 than the Stark profile of L_{β} , and broader by a factor 1.7 for H_{α} .



Figure 3. Same as Figure 1 for H_{α} .



Figure 4. Line shape of L_{α} for soliton only with W = 1.1 (solid line), for Stark only in the impact approximation (dashed line), and compared to a convolution (dotted line) of the latter two for the plasma condition of Figure 1.



Figure 5. Same as Figure 4 for L_{β} .



Figure 6. Same as Figure 4 for H_{α} .

5. Conclusions

The study of Zhakarov equations and numerous numerical simulations has revealed the complex behavior of plasma affected by strong Langmuir turbulence. The nonlinear coupling of the plasma waves creates numerous localized wave packets, subject to collapse. Each wave packet experiences a cycle during which the electric field magnitude grows to values of more than hundred E_0 ($E_0 \approx$ Holtsmark field). Using the main properties of strong Langmuir turbulence obtained from simulation calculations, we proposed a simple stochastic renewal model for the electric field of the wave packets. This model field is well-suited to study the effect of Langmuir turbulence on the line shape emitted in plasma. We used a simulation to generate field histories with a prescribed probability density function and waiting time distribution. The dipole autocorrelation function was obtained by a numerical integration of the Schrödinger equation and an average over 10⁴ histories. Our calculations concern the hydrogen L_{α}, L_{β}, and H_{α} lines for plasma conditions for a density equal to 10¹⁹ m⁻³ and a temperature of 10⁵ K. Strong turbulence brings a significant additional broadening to the pure Stark profile for all three lines.

In the future, we will look for other lines, plasma conditions, and wave collapse conditions, and make comparisons with line shapes observed in turbulent plasma.

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Article Doppler Broadening of Spectral Line Shapes in Relativistic Plasmas

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Abstract: In this work, we report some relativistic effects on the spectral line broadening. In particular, we give a new Doppler broadening in extra hot plasmas that takes into account the possible high velocity of the emitters. This suggests the use of an appropriate distribution of the velocities for the emitters. Indeed, the Juttner-Maxwell distribution of the velocities is more adequate for relativistic velocities of the emitters when the latter are in plasma with an extra high temperature. We find an asymmetry in the Doppler line shapes unlike the case of the traditional Doppler effect.

Keywords: plasmas; Maxwell; Juttner-Maxwell; relativistic; Doppler effect; asymmetry

1. Introduction

The Doppler effect, discovered by physicist and mathematician Christian Doppler in the nineteenth century, is the modification of the frequency of a wave when the emitting source and the receiver are in relative motion. The frequency change also implies that of the period and the wavelength. This effect concerns both mechanical waves and electromagnetic waves. In plasmas, the neutral atoms, molecules or ions moving inside the plasma are similar to the moving antennae. Atoms or ions subjected to the Doppler effect, exhibit the well-known phenomenon: the Doppler broadening of the line profile. The investigation in recent decades of the derivation and illustration of the Doppler effect, especially the generalized relativistic Doppler effect, is still being actively pursued today [1–5]. In the following we will illustrate this method to formulate the classical Doppler effect first, and then the same method is formulated to get the relativistic Doppler effect on the Doppler broadening of the line profile. In our work, we will present a better derivation allowing quick and exact expressions of the classic and relativistic Doppler effect on the broadening of the spectral lines observed in the plasmas. In the formulation of the classical Doppler effect, we used the Maxwell velocity distribution for the emitters, while in the formulation of the relativistic Doppler effect we used the relativistic Juttner-Maxwell distribution. The latter is justified for the case of very high temperatures (in the range 10⁵–10⁸ K) such as that encountered in fusion plasmas, in astrophysics, in cosmology (primordial Universe) and in unstable Z Pinch experiments [5]. Indeed, the Juttner-Maxwell distribution remains valid for all temperatures since it is more general than the Maxwell distribution.

2. Doppler Broadening

2.1. Classical Doppler Broadening: Non Relativistic Case

Often the emission (or absorption) of radiation by a particle (atom, ion, etc.) occurs during the movement. By the Doppler effect, the observed frequency in the observer (at rest) frame (see Figure 1) is different from the frequency emitted in the atom frame. The mean particle velocity at thermodynamic equilibrium is related to the temperature of the medium. Hence the broadening of the statistical Doppler effect is related to the distribution of the velocities of the emitter at the temperature T of the medium and the mass m of the emitter.



Figure 1. The fixed frame where the emitter moves with a velocity *V* forming an angle θ with the observation direction Ox.

One can assume a motionless observer, looking at an emitting atom moving with a velocity V in a direction forming an angle θ with the direction of observation (Ox) (see Figure 1), records a shifted angular frequency ω with respect to the angular eigenfrequency ω_0 of the emitter assumed to be stationary. This angular frequency is given by

$$\omega(V_x) = \omega_0(1 - \frac{V}{c}\cos\theta) = \omega_0(1 - \frac{V_x}{c})$$
(1)

where *c* is the velocity of the light in vacuum. The normalized intensity (normalized to one) of the line at the angular frequency ω is given by the average over the normalized Maxwell distribution (normalized to one).

$$f_{Maxwell}(V_x) = (m/(2\pi k_B T))^{1/2} \exp(-mV_x^2/(2k_B T))$$
(2)

of the Dirac delta distribution as the following

$$I(\omega) = \langle \delta(\omega - \omega(V_x)) \rangle_{Maxwell} = \sqrt{\frac{m}{2\pi k_B T}} \int_{-\infty}^{+\infty} \exp(-\frac{m}{2k_B T} V_x^2) \delta(\omega - \omega(V_x)) dV_x$$
(3)

Using the integral representation of the Dirac delta distribution (u is the integration variable whose unit is the second)

$$\delta(\omega - \omega(V_x)) = \frac{1}{2\pi} \int_{-\infty}^{+\infty} \exp(iu(\omega - \omega(V_x))) du$$
(4)

We find the normalized intensity (normalized to one) as

$$I(\omega) = \frac{1}{2\pi} \sqrt{\frac{m}{2\pi k_B T}} \int_{-\infty}^{+\infty} \exp(iu\omega) du \int_{-\infty}^{+\infty} \exp(-\frac{m}{2k_B T} V_x^2 - iu\omega(V_x)) dV_x$$
(5)

$$=\frac{1}{2\pi}\sqrt{\frac{2\pi mc^2}{k_B T \omega_0^2}}\exp\left(-\frac{mc^2}{2k_B T}(\widehat{\omega}-1)^2\right)$$
(6)

where $\hat{\omega} = \frac{\omega}{\omega_0}$, k_B is the Boltzmann constant and *m* is the emitter mass. This is the formula of the intensity of the line in the non relativistic case. It is symmetric (Gaussian) around the central angular frequency ω_0 . We note that the integrals in Formula (6) are strongly convergent because we deal with purely Gaussian integrals. The full width at the half maximum (FWHM) is given by the well known formula (in angular frequency unit)

$$\Delta\omega_{Doppler} = \omega_0 \sqrt{\left(\frac{8k_B T \ln(2)}{mc^2}\right)} = 7.1574 \times 10^{-7} \times \omega_0 \sqrt{\frac{T}{M}}$$
(7)

where M is the mass of the emitter in atomic mass unit whereas T is the temperature in Kelvin.

2.2. Relativistic Doppler Broadening

When an observer at a rest, recording the emitted radiation from a moving atom (or ion) with relativistic velocity *V*, they find that the angular frequency of this radiation is equal to [6]:

$$\omega(\beta) = \omega_0 \gamma (1 + \beta \cos \theta) \tag{8}$$

where

$$\beta = V/c, \tag{9}$$

$$\gamma = 1/\sqrt{(1-\beta^2)} \tag{10}$$

and ω_0 is the angular eigenfrequency and θ is the angle between the velocity of the emitter and the observation direction (*Ox*) (see Figure 1). By using the normalized Juttner-Maxwell distribution (normalized to one) [7]

$$W_{J-M}(\beta)d\beta = \lambda \frac{\gamma^5 \beta^2 d\beta}{K_2(\lambda)} \exp\left(-\lambda\gamma\right) \tag{11}$$

where

$$\lambda = mc^2/(k_B T) \tag{12}$$

and $K_2(X)$ is the modified Bessel function of order two, we obtained the normalized relativistic intensity (normalized to one) of the line profile

$$I(\omega) = \langle \delta(\omega - \omega(\beta)) \rangle_{Juttner-Maxwell} = \int W_{J-M}(\beta) d\beta \cdot \delta(\omega - \omega(\beta))$$
(13)

$$=\frac{1}{4\pi}\int_{-\infty}^{+\infty} du \int \int \int W_{J-M}(\beta) d\beta \exp(iu(\omega-\omega(\beta)))\sin\theta d\theta d\phi$$
(14)

Here, we have introduced the integral over the spherical angles that makes the emitter velocity with the fixed frame axis (see Figure 1). We have replaced the Dirac delta distribution by its integral representation by integrating over the variable *u*. Replacing the Juttner-Maxwell distribution $W_{J-M}(\beta)$

given by (11) and $\omega(\beta)$ given by (8) in Formula (14), we reach a more suitable expression of the relativistic intensity of the line profile

$$I(\omega) = \frac{\lambda}{4\pi \cdot K_2(\lambda)} \int_{-\infty}^{+\infty} \exp(iu\omega) du \int_0^1 \gamma^5 \beta^2 \exp(-\lambda\gamma) \exp(-iu\omega_0\gamma) d\beta \int_0^{\pi} \exp(iu\omega_0\gamma\beta\cos\theta) \sin\theta d\theta$$
(15)

or after integration on θ between zero and π ;

$$I(\omega) = \frac{\lambda}{2 \cdot K_2(\lambda)} \int_0^1 \gamma^4 \beta \exp(-\lambda\gamma) d\beta \int_{-\infty}^{+\infty} \frac{\exp(iu\omega - iu\omega_0\gamma + iu\omega_0\gamma\beta)}{iu\omega_0} du -\frac{\lambda}{2 \cdot K_2(\lambda)} \int_0^1 \gamma^4 \beta \exp(-\lambda\gamma) d\beta \int_{-\infty}^{+\infty} \frac{\exp(iu\omega - iu\omega_0\gamma - iu\omega_0\gamma\beta)}{iu\omega_0} du$$
(16)

Finally, the integration over *u*, allows us to get the relativistic intensity of the line profile

$$I(\hat{\omega}) = \frac{\lambda}{2 \cdot K_2(\lambda)} \int_1^\infty \gamma d\gamma \exp(-\lambda\gamma) \left(S(\hat{\omega} - \gamma + \sqrt{\gamma^2 - 1}) - S(\hat{\omega} - \gamma - \sqrt{\gamma^2 - 1}) \right)$$
(17)

where S(t) = +1 if t > 0 and S(t) = -1 if t < 0 and $\hat{\omega} = \frac{\omega}{\omega_0}$ is the reduced angular frequency. We can manage the formula to be more suitable for the numerical treatment:

$$I(\widehat{\omega}) = \frac{\lambda}{2 \cdot K_2(\lambda)} \exp(-\lambda) \int_1^\infty \gamma d\gamma \exp(-\lambda(\gamma - 1)) \left(S(\widehat{\omega} - \gamma + \sqrt{\gamma^2 - 1}) - S(\widehat{\omega} - \gamma - \sqrt{\gamma^2 - 1}) \right)$$
(18)

if we put $\lambda(\gamma - 1) = y$, (λ is given by formula (12)) we find

$$I(\widehat{\omega}) = \frac{\exp(-\lambda)}{2\cdot k_2(\lambda)} \int_0^\infty (\frac{y}{\lambda} + 1) dy \exp(-y) * \left(S(\widehat{\omega} - \frac{y}{\lambda} - 1 + \sqrt{(\frac{y}{\lambda})^2 + 2\frac{y}{\lambda}}) - S(\widehat{\omega} - \frac{y}{\lambda} - 1 - \sqrt{(\frac{y}{\lambda})^2 + 2\frac{y}{\lambda}})\right)$$
(19)

We note that the integral in the last formula is convergent because we deal with the integral in distribution sense [8].

Unlike the classical Doppler effect, the relativistic one has a property: an asymmetric broadening as it is shown clearly in the following table corresponding to the temperatures in the range 10^5-10^9 K. We remark also that in the relativistic case, as in the classical case, the central frequency is unchanged (see Figure 2). The maximum value of the intensity is at $\hat{\omega} = 1$ both for classical and relativistic case but the maximum value of the relativistic case is smaller than the maximum of the classical case (the asymmetry at $\hat{\omega} = 1$) is negative, see Figure 3.

We mention that in the Table 1, we have denoted by $\hat{\omega}_{L,R}$ the value of the reduced angular frequency at the left and the right of the peak of the line (centred at $\hat{\omega} = 1$). $\hat{\omega}_L$ and $\hat{\omega}_R$ are chosen to be symmetrical with respect the centre of the line at $\hat{\omega} = 1$ and giving intensities very close to the half of the maximum of the intensity. Strictly speaking, we have considered $\hat{\omega}_R = 1 + g$ and $\hat{\omega}_R = 1 - g$ with g = 0.00045 for Fe^{+25} , 0.00075 for W^{+73} , 0.00085 for Fm^{+99} (produced in nuclear reactions) and 0.00095 for Cn^{+111} (it is synthesized in laboratories for use in nuclear reactions). As we see in this table, the value of the intensity at the right $I(\hat{\omega}_R)$ is greater than the intensity at the left $I(\hat{\omega}_L)$. This remark shows clearly that the line profile has an asymmetry as defined by [9] (see the definition at the last line in the above table). Another feature in this study is that we have not specified the line profile, because we have used the reduced angular frequency $\hat{\omega} = \frac{\omega}{\omega_0}$: for each specific line (specific transition), we must multiply $\hat{\omega}$ (the x-axis) by the corresponding angular eigenfrequency ω_0 to obtain the intensity $I(\omega)$. If we define the asymmetry as [10]

$$Asym = I(\omega, relativistic) - I(\omega, classical)$$
(20)

which is the difference between the normalized relativistic intensity given by (18) and the normalized classical intensity given by (6), we obtain the following figure representing the asymmetry for Cn^{+111} at 1.9×10^9 K.



Figure 2. Relativistic intensity as defined by Formula (19) for Iron at $T = 10^{10}$ K.



Figure 3. Asymmetry as defined by Formula (20) for Cn^{+111} at T = 1.9×10^9 K.

Table 1. Asymmetry percentages for different hydrogen-like ions.

	$T = 10^8 K$, Fe^{+25}	T = 8.5 \times 10 8 K, W^{+73}	T = 1.5 \times 10 ⁹ K, Fm^{+99}	\mathbf{T} = 1.9 \times 10 9 K, \mathbf{Cn}^{+111}
$\hat{\omega}_L$	0.99955	0.99925	0.99915	0.99905
$\hat{\omega}_R$	1.00045	1.00075	1.00085	1.00095
$I(\hat{\omega}_L)$	1.0802	1. 0303	1.0195	0.95715
$I(\hat{\omega}_R)$	1.0807	1.0313	1. 0206	0.95849
$\frac{I(\hat{\omega}_R) - I(\hat{\omega}_L)}{I(\hat{\omega}_R) + I(\hat{\omega}_L)} * 100$	0.030	0.051	0.056	0.060

In Figure 3, we see clearly that, in the left of $\hat{\omega} = \omega/\omega_0 = 1.00007$, the intensity of the relativistic profile is lower than of the classical profile, whereas it is higher in the right of $\hat{\omega} = \omega/\omega_0 = 1.00007$. It can be seen clearly in this figure that the asymmetry is a function of $(\hat{\omega} = \omega/\omega_0)$ and that means that for any line, the asymmetry is as indicated in this figure. To obtain the asymmetry, for a specific line centred at w_0 , we must multiply $\hat{\omega}$ by w_0 . The same remark holds for Figure 4 for the hydrogen-like Iron, but with a more pronounced asymmetry since the maximum of the asymmetry is equal to 2.40 for the iron (Fe⁺²⁵) whereas for the Copernicium (Cn⁺¹¹¹) is equal to 2.12.



Figure 4. Asymmetry as defined by Formula (20) for Fe^{+25} at T = 1.9 × 10⁹ K.

3. Conclusions

In this work, we report some relativistic effects on the spectral line broadening. In particular, we obtained a new expression for the Doppler broadening that takes into account the possible high velocity of the emitters. This suggests the use of an appropriate distribution of the velocities for emitters. We find, an asymmetry in the Doppler broadening unlike the well known classical Gaussian Doppler broadening.

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Article Active Galactic Nuclei Search

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Abstract: We present the first results of medium-band photometric observations on the 1m Schmidt Telescope of Byurakan Astrophysical Observatory (Armenia). The object sample was created in the SA68 field. The medium-band filter set (13 filters with FWHM = 250 Å + 5 broadband SDSS filters) allowed us to create low-resolution spectra of each object in the SA68 field. We compared them with the template spectra to select AGNs and to determine their photometric redshifts. Our sample consists of 330 objects with 0.5–5.1 redshift range and complete up to 23.0 AB magnitude. The comparison of our sample with SDSS DR10 and BOSS + MMT QSO showed that sufficiently more objects in the 3.2–5.1 redshift range were found.

Keywords: QSO samples; QSO selection; medium-band photometry

1. Introduction

In studies of quasar evolution we need to create a homogeneous sample of objects. The wide variety of methods to obtain such samples exists, and the most commonly used ones are described below.

1.1. Radio Surveys

Radio surveys are very powerful for searching AGN and were the original method of QSO selection. It works well at high luminosities, since almost all luminous radio sources are active nuclei. Additionally, the significant part of radio galaxies shows no strong evidence of non-thermal activity [1]. On the other hand, only near 10% of all AGN are "radio loud," so the radio samples are respectively incomplete. Furthermore, the ratio of bolometric luminosity to radio luminosity varies by a factor of 10⁵, and the fraction of the total energy radiated in the radio is small [1].

1.2. X-ray Surveys

The X-ray advantage over the optical selection is that stars do not have high luminosities in this range; consequently, it greatly simplifies the selection of objects. However, to achieve sufficient depth for the detection of distant quasars, long exposure times are required, which is possible only in small fields. Thus, the samples consist of a small object number, and statistical studies based on them are not reliable (Figure 1) [2]. Additionally, the brightness of the quasars at the X-ray range strongly depends on the orientation of the active nucleus, that the part of the active nuclei in X-rays are faint for detection.



Figure 1. The number density of quasars from X-ray data [2]. Solid, dashed, and dash-dotted lines show AGN number densities for different luminosity ranges.

1.3. Broadband Optical Photometry in the Combination with Infrared and Ultraviolet Surveys

Broadband photometry as a method of active nuclei search has been used for approximately 50 years [3]. With a small number of broadband filters, observers can reach an appreciable depth with short exposure time and can classify field objects based on their color excesses. However, for quasar searching, this technique is confined: shifted emission lines, appearing in different filters, generate a large spread of values in the color space, and the ultraviolet excess does not reveal itself at z > 2.2. One possible solution is to supplement optical photometry with the data from infrared and ultraviolet surveys [4]. Yet, another difficulty is faced: one of the main selection criteria for the WISE infrared data (W1 – W2 > 0.8 proposed by [4]) stops working properly when z > 3 because a significant fraction of quasars is below this limit, which affects sample completeness (Figure 2) [5]. At large redshifts (z > 3), the Lyman break appears at the optical range, which theoretically makes it possible to separate quasars from stars. However, in practical terms, the color difference between two nearby broadband filters should be greater than 2^m. Thus, for SDSS, at a magnitude limit of about 22^m, we can reliably see this difference for objects no weaker than 20^m, which allows us to observe only the brightest quasars at large redshifts. Active nuclei recorded with a lower signal-to-noise ratio are indistinguishable from red stars.



Figure 2. The WISE infrared colors of AGNs. Central line shows averaged color-redshift diagram for selected AGNs; upper and lower lines shows data errors. It can be clearly seen that, for most quasars at z > 3, the W1 – W2 > 0.8 criterion is not met [5].

1.4. QSO Selection by Variability in the Optical Range

Most AGNs are variable objects at the long-periodical timescale and can be used as a selection criterion. This was implemented by Palanque-Delabrouille et al. using E-BOSS data [6]. Theoretically, quasar selection by variability does not require the use of colors. Practically, however, with only variability selection, some variable stars were included in the sample, and additional color selection was still essential (Figure 3) [6].



Figure 3. The BOSS + MMT color selection criteria [6]. Locus of stars (upper blue contours), z < 2.2 quasars (lower left green contours), and z > 2.2 quasars (lower right red contours) in the c3 vs. c1 color-color plane. The upper solid line corresponds to the color cut c3 < 1.0 - c1/3 (loose, for point sources) and the lower dashed line to c3 < 0.6 - c1/3 (strict, for extended sources). c1 = 0.95(u - g) + 0.31(g - r) + 0.11(r - i), c3 = -0.39(u - g) + 0.79(g - r) + 0.47(r - i), where u, g, r, i and z are SDSS colors.

1.5. Medium-Band Photometry

One of the problem solutions for quasars selection in the optical range is to increase the spectral resolution. By increasing the number of filters and by decreasing their width, we can obtain a spectral energy distribution of each of the field objects with a higher resolution, while the exposure time required to reach the same depth increases as well. Such an approach makes it possible to detect broad emission lines in the AGN spectrum, which presents an opportunity both to separate quasars from stars and to determine their photometric redshifts with sufficient accuracy (Figure 4). This method of AGN selection is self-sufficient, but the addition of medium-band photometry with infrared and X-ray data can greatly simplify the selection.

We examined quasar samples obtained with three different methods: broadband optical photometry in combination with infrared and ultraviolet data (SDSS DR10, [7]), optical variability selection (BOSS + MMT, [6]) and X-ray data ([2,8]). At large redshifts (z > 3), there is a decrease in the number of objects from the SDSS DR10 broadband photometry (Paris et al., 2013) and the BOSS + MMT sample ([6]). However, according to X-ray data, there is no such noticeable decline ([2,8]). Based on the assumption that the lack of distant active nuclei in optical surveys is due to selection effects, we decided to make a QSO sample using medium-band photometry and started the "Medium-Band Byurakan Survey" (MBBS).



Figure 4. The accuracy of the photometric redshift determination from the medium-band photometry data. Along the vertical axis—the photometric redshift of quasars, along the horizontal—their spectroscopic redshift. $\sigma = 0.02$ [9].

2. Observations

The medium-band survey of the SA68 field was made on the 1-m Schmidt telescope at the Byurakan Astrophysical Observatory of the National Academy of Sciences of Armenia (BAO). In 2013–2015 the Laboratory of Spectroscopy and Photometry of Extragalactic Objects of the Special Astrophysical Observatory of Russian Academy of Sciences together with the Armenian specialists was engaged in its modernization. The 1-m telescope of the Schmidt system (105/132/213) of the BAO is one of the largest instruments in the world: it is one of the five largest telescopes in terms of mirror size and the third in terms of the objective prisms size. In October 2015, we installed a CCD detector (4 k × 4 k, liquid-cooled, RON ~11.1 e, manufactured by Apogee, Santa Monica, CA, USA) in the telescope focus with a resolution element of 0.868 arcsec, a field of view of about 1 square degree. The detector is equipped with filter wheels with 20 medium-band filters (FWHM = 250 Å, spectral range covering 4000–9000 Å), 5 broadband filters (u, g, r, i, z SDSS), and 3 narrowband filters (5000 Å, 6560 Å and 6760 Å, FWHM = 100 Å) (Figure 5). Observations were made by the authors of this paper over a period of two weeks in October and November 2016. Such a small observation period minimizes the influence of long-period AGN variability in the observational data.



Figure 5. Transmission of broadband (blue lines) and medium-band filters (black lines), taking into account the quantum efficiency of the detector (red dot-dash line). With the dash line, we illustrate a typical night sky spectrum. Filters were measured at laboratory with F/2 beam, corresponding to the focal ratio of the 1-m telescope.

3. Photometry and SED Construction

The SA68 field was exposed in 13 medium-band and 5 broadband filters. The width of medium-band filters was chosen so that the spectral resolution was sufficient to register the broad emission lines (Figure 6). To avoid color selection problems during the object detection, a basic object sample was created with a deep image, taken by summing g, r, and i filters (the limiting magnitude in broadband filters was about 25^{m}). The sample was cut to 23^{m} , which is explained by the depth of the medium-band images, for which exposure times of 40-60 min allowed us to register up to 23^{m} objects with a signal-to-noise ratio higher than 5. This allow us to create the luminosity function of the quasars with MB > -23 up to Z = 3.2 and up to Z = 5 for objects with MB > -24.7. The master list was then created based on deep-image photometry (Figure 7). Thus, medium-band photometry of all field objects was performed, and spectral energy distribution [9] was constructed (Figure 7).



Figure 6. Some of medium-band selected AGN characteristics. The left images shows the object positions. The center images show medium-band SEDs (black dots with solid line) and broad-band SEDs (triangles with dashed line). The spectra of objects obtained with the 6-m telescope of SAO RAS are shown on the right.



Medium Band Byurakan Survey

Figure 7. Cont.



Figure 7. (a) Photometry algorithm scheme and (b) scheme of SED construction based on medium-band data [9].

4. QSO Selection

The primary selection of quasar candidates was made automatically by comparing the SED of objects with the SED calculated from the model template spectra of the active nuclei. The probability that the object is active galactic nuclei was calculated, and the photometrical redshift determined, using LePhare templates package ([10,11]). Then, the SEDs of the selected objects were viewed manually to make sure the selection was correct.

5. Results

According to the selected objects, the dependencies of the quasar number density on the redshift were constructed, and a direct comparison was made with the SDSS DR10 and BOSS + MMT samples. It can be clearly seen that, for BOSS + MMT, a significant decrease in spatial density begins at z > 2.2; for SDSS DR10, a similar decline is noticeable at z > 3.5 (Figure 8). However, according to the MBBS data, there is no such significant decrease in object number, which is in good agreement with the X-ray data.



Figure 8. The number densities of quasars based on three different samples. $\Delta z = 0.2$. The green line shows the QSO number density based on the SDSS data release 10 sample [7], and the red line shows that based on BOSS + MMT variability selection sample [6]. The black line shows the number density based on the Medium-Band Byurakan Survey sample in the SA68 field.

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Article

Models of Emission-Line Profiles and Spectral Energy Distributions to Characterize the Multi-Frequency Properties of Active Galactic Nuclei

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Abstract: The spectra of active galactic nuclei (AGNs) are often characterized by a wealth of emission lines with different profiles and intensity ratios that lead to a complicated classification. Their electromagnetic radiation spans more than 10 orders of magnitude in frequency. In spite of the differences between various classes, the origin of their activity is attributed to a combination of emitting components, surrounding an accreting supermassive black hole (SMBH), in the *unified model*. Currently, the execution of sky surveys, with instruments operating at various frequencies, provides the possibility to detect and to investigate the properties of AGNs on very large statistical samples. As a result of the spectroscopic surveys that allow the investigation of AGNs. In this contribution, we present the results obtained by working on multi-frequency data, and we discuss their relations with the available optical spectra. We compare our findings with the AGN unified model predictions, and we present a revised technique to select AGNs of different types from other line-emitting objects. We discuss the multi-frequency properties in terms of the innermost structures of the sources.

Keywords: galaxies: active; galaxies: nuclei; line: profiles; quasars: emission lines; catalogs; surveys

1. Introduction

When discussing active galactic nuclei (AGNs), we generally refer to the nuclei of galaxies for which a supermassive black hole (SMBH), with a mass ranging between 10^6 and $10^{10} M_{\odot}$, is fed by a continuous flow of matter from the surrounding environment. This process, denoted as *accretion*, leads to the conversion of the gravitational binding energy of the accreted material into heat and radiative energy, through the effects of the viscous interactions that arise in the accreted matter as it is accelerated up to several thousand kilometers per second by the strong gravitational pull of the black hole [1]. In spite of this common interpretation, AGNs present a wide range of striking observational differences in their spectra, in their total power and in the frequency range for which most of their energy is radiated away. While the first to be clearly identified were located in galaxies with exceptionally bright optical nuclei [2], nearly 10% of the total population were subsequently found to radiate large fractions of their power in the radio and the high-energy domains [3]. Their total luminosities can change over

a wide range and are typically considered to lie between $10^{40} \text{ erg s}^{-1}$, in the case of low-luminosity sources, and some $10^{46} \text{ erg s}^{-1}$ for the most powerful sources. They are distributed from the local universe, where the low-luminosity objects are more common, all the way up to very high redshifts ($z \ge 7$), where powerful activity becomes more frequent.

In terms of the characteristics of the optical-UV spectra, AGNs are generally characterized by the presence of a non-thermal continuum, often well represented by a power-law shaped spectrum of the form $L_{\nu} \propto \nu^{-\alpha}$ and sometimes accompanied by prominent emission lines with different profiles. In some cases, we additionally observe a hot thermal excess, with a peak that likely falls in the far UV, or different contributions from the host-galaxy stellar populations. Following a scheme that was first outlined by [4], we generally classify Type 1 AGNs as those objects whose spectra show broad recombination lines, with profiles corresponding to velocity fields exceeding 1000 km s⁻¹ in full width at half maximum (FWHM), from H, He, or from other permitted lines of heavy ions such as Fe II, C IV and Si IV, together with narrow forbidden lines (FWHM \approx 300–500 km s⁻¹) from, for example, [Ne V], [Ne III], [O II], [O II], [N II] and [S II]. We classify Type 2 AGNs as those for which both permitted and forbidden lines only have narrow profiles. In general, it is observed that Type 1 sources are brighter and commonly show a thermal UV excess, while Type 2 objects are dimmer and more severely contaminated by the host-galaxy spectral contributions.

The most widely accepted way to explain the observations is to assume that the central accreting SMBH is surrounded by a hot accretion disk, radiating in the optical, UV and X-ray frequencies, and a compact region (less than 0.1 pc in size) of dense ionized gas ($N_e \ge 10^9$ cm⁻³), which produces Doppler-broadened recombination lines due to the large gravitational acceleration, and is therefore called the *broad-line region* (BLR). The gas that is located at larger distances (1 pc $\leq r \leq 1$ kpc); although still being ionized and producing emission lines, it has a considerably smaller velocity field and electron densities closer to typical nebular environments ($10^3 \,\mathrm{cm}^{-3} \le N_{\ell} \le 10^6 \,\mathrm{cm}^{-3}$). It, therefore, radiates both permitted and forbidden lines with narrow Doppler profiles, giving rise to what we call the narrow-line region (NLR). If the central structure is partially obscured by an optically thick distribution of matter, as is supported by some observational evidence [5], the difference between Type 1 and Type 2 objects is consistently explained by the fact that our line of sight, respectively, may or may not reach the central regions, without being intercepted by the surrounding material, in what is called the AGN unified model [6]. When the accretion flow becomes coupled with the magnetic fields that develop close to the SMBH in such a way that a relativistically beamed jet of plasma is accelerated away from the nucleus, AGNs become powerful sources of radio and high-energy emission, eventually developing extended radio morphologies [7].

In spite of the fairly comprehensive interpretation, AGNs still pose many fundamental questions, because most of the relevant physical processes involved in the accretion and in the acceleration of jets are confined close to the SMBH, in a region that lies still beyond the resolution capabilities of present-day instruments. For this reason, a large part of our current knowledge concerning AGNs is based on the analysis of their spectra and on monitoring the correlations existing among their numerous spectral components. However, not all sources are equally good for such investigations, as Type 1 objects tend to be dominated by the emission of the AGN, while Type 2 sources are strongly affected by obscuration and stellar light contributions from the host galaxy. The main details to understand the physics of AGNs, therefore, can only be constrained by careful inspection of spectral properties that should be possibly extended over large statistical samples, in order to compare the predictions of different models with the available observations. In recent times, several campaigns have been carried out to monitor the sky at different frequencies and to obtain spectroscopic observations. In this contribution, we describe a revised scheme to select AGNs on the basis of the properties of their emission lines and colors, we illustrate the potential of multi-frequency models to relate the observed spectral energy distributions (SEDs) with optical spectra, and we discuss the information that we are able to extract on their central structures from the combined analysis of line profiles and multi-frequency data.

2. Results

The most common property shared by different types of AGNs is the emission of an intense continuum of non-thermal radiation that can possibly ionize diffuse gas and, thus, give rise to emission lines in the spectra. Because lines excited by non-thermal ionizing continua differ in intensity and distribution from lines excited by the continuum of hot thermal sources [8,9], we can apply a set of diagnostic diagrams, based on the intensity ratios of specific lines, in order to recognize the footprint of ionization from thermal and non-thermal sources in external galaxies. When the amount of spectroscopic data was dramatically increased, as a result of the execution of large spectroscopic surveys such as the *Sloan Digital Sky Survey* (*SDSS*) [10,11], this method was further refined, demonstrating its ability to recognize the presence of obscured AGN activity [12]. This kind of information is fundamental to assess the statistical relations existing between obscured and unobscured sources, which constrain the structure of the central source and its possible dependence on luminosity or age. In order to perform such a study, however, we need an instrument that is in principle able to detect different types of AGN activity, with the smallest possible influence of selection effects. We obtained such a tool by combining spectroscopic and photometric parameters, on the basis of an investigation of how different spectral classes are related with multi-frequency emission.

2.1. AGN Selection from Spectroscopic Surveys

The investigation of AGN statistical properties requires the revising of the selection techniques, which have been classically designed to detect specific types of sources on the basis of their characteristic properties. While Type 1 AGNs are generally well identified by the presence of prominent broad emission lines in the spectra, which are commonly accompanied by a hot thermal continuum excess sometimes referred to as the *big blue bump*, Type 2 AGNs are only characterized by narrow emission lines, which can also be present in the spectra of galaxies with strong star formation activity. In the case of narrow line-emitting sources, the methods based on diagnostic diagrams are fairly well suited to distinguish between AGN and stellar activity, but if performed on a selection of spectra simply based on the presence of emission lines, to collect different types of sources, the emission-line diagnostics alone are not straightforwardly applicable. An example of this effect is shown in Figure 1, where we compare different methods to distinguish AGNs from thermally excited spectral line emitters. The reason that the classic diagnostic ratios cannot be used on a sample of objects including Type 1 sources is attributed to the use of the recombination lines, which are needed to normalize the strength of the forbidden lines that probe the temperature and the ionization structure of the gas. The presence of a strong contribution in the broad component of the recombination lines in Type 1 objects, which is not balanced in the forbidden lines, forces Type 1 sources to populate the non-AGN region of the plots. This is apparent, because this type of diagnostic diagram is designed to work on the emission of the NLR alone and cannot account for the BLR component. It has been proposed that a different choice of the emission-line diagnostic ratios, involving only forbidden lines, might in principle solve the problem [13], but the available choices either involve the use of weak lines, or they are strongly subject to the effects of interstellar extinction.

If the direct sight of the central regions in Type 1 AGNs can give rise to problems in recognizing their spectral signature, on the other hand, we have the opportunity to take advantage of the strong blue and UV continuum, which is produced by the central source and is not obscured along the line of sight. It has been shown that Type 1 AGNs can effectively be selected by means of photometric criteria that compare their colors with those of non-active objects, to the extent that the SDSS uses a photometric pipeline to select QUASAR candidates for follow-up spectroscopy [14,15]. This method has some limitations when comparing objects at very different redshifts; however, in the low-redshift domain in which the diagnostic lines are still available in the optical frequency window ($z \le 0.5$), it defines a well-established parameter space in which Type 1 AGNs can be effectively distinguished from other line-emitting sources. A projection of this parameter space on the u - r versus g - z color–color diagram is also illustrated in Figure 1. The choice of these extended color bands, which

was based on the extinction-corrected magnitude measurements of all SDSS objects with emission lines of H α , H β , [O III] λ 5007 and [N II] λ 6583 detected at more than the 5 σ level, maximizes the effect of the blue continuum of Type 1 sources over the stellar continuum of other sources. As a consequence, we recover the possibility of detecting different types of nuclear activity by combining classic spectroscopic diagnostics and photometric colors into a multi-dimensional parameter space, in which AGNs populate a separate sequence with respect to other non-AGN powered sources.



Figure 1. Distribution of active galactic nuclei (AGNs; blue points) and star formation (red points) powered objects with emission-line spectra on the [O III] λ 5007/H β vs. [N II] λ 6583/H α diagnostic diagram (**left panel**), on the u - r vs g - z color–color diagram (**middle panel**) and on the three-dimensional parameter space combining the two-color photometry with the [O III] λ 5007/H β diagnostic ratio. The distribution in the three-dimensional parameter space illustrates how Type 1 AGNs are best identified among line-emitting sources by means of photometric criteria, while Type 2 sources can be distinguished on the basis of their spectral properties.

2.2. Emission Lines and Models of AGN Spectral Energy Distributions

The selection of general samples of AGNs belonging to different spectral classes allows us to search for observations of the corresponding sources in multiple frequencies. By combining the available data, it is possible to reconstruct the AGN SEDs and to compare these with the associated optical spectra, as is illustrated for example in Figure 2. The plots show the different SEDs of two prototypical AGNs (3C 273 for Type 1 and NGC 1068 for Type 2) modeled through a combination of thermal and non-thermal radiation components, together with their characteristic spectra. We can immediately appreciate how the occurrence of a Type 1 spectrum, with broad lines and a blue continuum, is well associated with a strong dominance of the non-thermal contribution and a direct sight towards the hottest central regions, resulting in an excess of ionizing radiation, in agreement with the unified model predictions. Conversely, the Type 2 SED is totally consistent with an obscured central source, whose low-energy ionizing radiation is severely absorbed and reprocessed by a distribution of material that subsequently re-emits photons in the infrared (IR) domain. Only the more penetrating high-energy photons and the long-wavelength radio emission, which is practically unaffected by obscuration, can propagate directly from the central source, therefore resulting in an optical spectrum that is dominated by the host galaxy and the emission lines originating from the unobscured NLR.

In addition to providing observational evidence in support of the unified model, and possibly to associate different degrees of absorption and light reprocessing to various AGN classes, the combination of SED models and spectroscopic observations is a promising instrument to improve our knowledge of AGN distribution. As a result of the numerous efforts that have been devoted to surveying the whole sky at various frequencies, it is now possible to recognize AGN candidates from their SED, even in sky areas that have not yet been covered by detailed and publicly available spectroscopic programs. The further possibility that models of the observed SED may lead to a prediction of the expected AGN class, in addition to optimizing the execution of follow-up campaigns, provides statistical constraints to infer structural and evolutionary details in large samples of AGNs.



Figure 2. Examples of spectral energy distribution (SED) models (**left panel**) and optical spectra (**right panel**) for the prototypical Type 1 active galactic nucleus (AGN) 3C 273 (blue line and points) and the prototypical Type 2 AGN NGC 1068 (red line and points). The SEDs have been normalized to the same optical V-band magnitude and the models are based on the combination of two cut-off power laws and one black-body contribution. The spectrum of 3C 273 was observed at the Asiago Astrophysical Observatory, while the spectrum of NGC 1068 is taken from literature [16]. Both spectra were taken to rest-frame wavelength scale.

2.3. Multi-Frequency Analysis of the Central Engine

Because of the extremely compact size of the regions in which the continuum and the bulk of the emission lines are produced in AGNs, we do not yet have a fully developed interpretation of their innermost structures. Most of our current understanding is derived from the analysis of spectra and from models that carry out the inferred physical conditions. Therefore, the extension of spectroscopic analysis to different frequencies improves our ability to explore unresolved structures. Such an example is illustrated in Figure 3 for the case of the QSO PG 1114 + 445, for which we compare the *XMM Newton* X-ray spectrum with measurements of the normalized intensities of the broad components of the Balmer lines of hydrogen. This quantity is defined as

$$I_n = \frac{\lambda_{ul} I_{ul}}{g_u A_{ul}} \tag{1}$$

where λ_{ul} is the wavelength corresponding to the transition from an upper level *u* to a lower level *l* (with l = 2 for H Balmer lines), I_{ul} is the measured intensity, g_u is the statistical weight of the upper level, and A_{ul} is the spontaneous transition probability. In the case of an optically thin line, we can use the general expression:

$$I_{ul} = \frac{1}{4\pi} \frac{hc}{\lambda_{ul}} A_{ul} \int_0^{s^*} N_u \mathrm{d}s \tag{2}$$

where *h* and *c* are the Planck constant and the vacuum speed of light, while N_u is the concentration of atoms in the upper level, and the integration is executed throughout the extension of the source to obtain

$$I_n = \frac{1}{4\pi} \frac{hc}{g_u} \int_0^{s^*} N_u \mathrm{d}s \tag{3}$$

If we now assume that the distribution of emitting atoms is spatially constant within the source and that it can be represented by a Boltzmann formula at least in the high-excitation stages (a condition known as *partial local thermodynamic equilibrium*—PLTE), we obtain Atoms 2017, 5, 43

$$I_n = \frac{1}{4\pi} \frac{hc}{g_1} N_1 s^* \exp\left(-\frac{\Delta E_u}{k_B T_e}\right) \tag{4}$$

where ΔE_u is the upper-level excitation energy. It is therefore clear that, under the assumed conditions, we expect the following:

$$\log I_n = \log \left(\frac{1}{4\pi} \frac{hc}{g_1} N_1 s^*\right) - \frac{\log e}{k_B T_e} \Delta E_u \tag{5}$$

which is a linear function of the upper-level excitation energy [17].



Figure 3. Suppression of high-order Balmer lines, observed through the normalized intensities of the broad components in the optical spectrum of PG 1114 + 445 (**left panel**), compared with the soft X-ray spectrum (**right panel**) obtained from the *XMM Newton* EPIC instrument (green points) and from the twin MOS cameras (red and black points). The X-ray spectrum is modeled with a power-law spectrum, a soft X-ray thermal excess and an ionized absorber. Absorption from the neutral medium within the Milky Way is also taken into account.

The deviation from the expected linear behavior observed in Figure 3 is an indication that the flux of high-order Balmer-line photons is lower than the prediction, which can happen in the presence of a dense layer of recombining plasma. In this case, indeed, Equation (2) would be modified in as

$$I_{ul} = \frac{1}{4\pi} \frac{hc}{\lambda_{ul}} A_{ul} \int_0^{s^*} N_u e^{-k_{lu}s} \mathrm{d}s \tag{6}$$

where

$$k_{lu} = N_l \sigma_{lu} \tag{7}$$

is the line absorption coefficient, controlled by the density of ions in the lower level N_l and the line photon absorption cross-section σ_{lu} . Under typical nebular conditions, this would be $k_{lu} << 1$ for any l > 1; however, in the presence of a thick layer of ionized plasma, such as that responsible for the observed absorption in the soft X-ray spectrum, which has an estimated column density of $N_C = 4.896 \times 10^{21}$ cm⁻², this could no longer be negligible. Under such circumstances, indeed, we expect the lower level of the Balmer series to be overpopulated by recombination processes and by Ly α photon trapping. If, therefore, this X-ray absorber is located outside the BLR, it could very likely also be responsible for the absorption of Balmer photons.

To further explore the possibility that the presence of ionized gas layers could affect the broad emission lines, we began an investigation into the Balmer-line profiles in the broad component of the spectra of a sample of narrow-line Seyfert 1 galaxies (NLS1s), selected from the SDSS Data Release 7 [18]. Because of the relatively small width of their broad-line components (FWHM_{H β} < 2000 km s⁻¹ [19]), the profiles of these lines are largely unaffected by blends with other broad lines, although the narrow components

are still to be accounted for. These, however, are much simpler to model, taking the narrow forbidden lines of [O III] as templates to constrain their widths and to be subtracted from the global profiles. By taking the broad-line components in a velocity scale, we are able to compare the resulting profiles, as illustrated in Figure 4. Using a narrow emission-line width fixed at 0.75 times the width of [O III] λ 5007, in order to account for the larger velocity dispersion of the high-ionization gas [20], we modeled the narrow components of H β , H α and the [N II] $\lambda\lambda$ 6548, 6583 doublet, attempting to isolate the broad emission-line profiles. Regarding the resulting FWHM of H α and H β , we find that most of the line profiles are very similar, favoring the interpretation of an optically thin gas, although we still observe deviations from this behaviour. It has already been noted that the profiles of these lines can be different in some objects and that they may even exhibit different reverberation lags [21,22]. This could point towards a displacement of their emission sites or to a relevant role of dust absorption in the central regions of the source [23,24], as there is convincing evidence that a substantial amount of dust can exist in the central regions of AGNs, at smaller scales than the NLR [25].



Figure 4. (Left panel) comparison of the H β (blue) and H α (red) broad line components, after subtraction of the narrow emission lines, plotted in velocity scale and normalized to the same peak intensity in the spectrum of 1RXS J113247.0 + 062626. (**Right panel**) distribution of the ratio FWHM(H α)/FWHM(H β) in the broad components of the emission lines from the spectra NLS1 galaxies investigated in [18]. While the distribution is strongly peaked around 1, favoring the interpretation of an optically thin gas, some differences in the line profiles can arise as a consequence of dust absorption or emission-line radiation transfer effects.

3. Discussion

The processes that occur in the unresolved central regions of AGNs leave characteristic signatures in the emission and absorption components of the observed radiation, and they also control which parts of the source are visible. Although a preliminary analysis of the relationships existing between multi-frequency data and optical spectra argue in favor of the unified model, a systematic study that collects the huge amount of available observational material still has to be carried out. This type of investigation is highly desirable because of the invaluable constraints that it could place on the nature and evolution of the innermost structures of AGNs, but it presents obvious difficulties related to the amount of data that should be considered. With our work, we present a technique to select different types of AGNs, with available spectro-photometric measurements, and we provide some examples of how the comparison of their spectra with the overall SED lead to supporting the unified picture, as well as to the intriguing possibility that SEDs built on the basis of public data could be used to select targets for follow-up spectroscopy or even to attempt preliminary classifications.

A particularly interesting result is that based on the combination of optical and X-ray spectroscopic observations. In the presence of a significant layer of ionized plasma, such as that observed in the X-ray spectrum of PG 1114 + 445, optical depth effects become important for determining the relative strengths of the recombination lines. In particular, these could explain the various emission-line intensity ratios that are known to deviate significantly from the standard recombination predictions in Type 1 sources. A substantial increase in optical depth gives rise to two important effects. On the one hand, the H α photon has a larger absorption cross-section with respect to higher-order Balmer photons. On the other hand, it corresponds to a transition between two adjacent levels and, therefore, has a smaller probability to decay through other spontaneous transition channels, with respect to higher-order Balmer photons. Therefore, an increase in the line optical depth might lead to a reduction of the high-order Balmer line intensities with respect to H α , but H α could averagely emerge from an outer layer of the line-emitting region. In a dynamical configuration dominated by Keplerian motions, this would imply a narrower line profile. On the other hand, the role played by dust in the emission-line regions would possibly lead to the opposite result, suppressing the short-wavelength, high-order emission lines more severely than $H\alpha$, therefore favoring an enhancement of this line from deeper regions. Comparing the measurements of the width of the broad emission-line profiles, as illustrated in Figure 4, shows that neither effect is likely to severely affect the emission in the vast majority of sources, although some exceptions may still exist. This result favors the possibility that the optically thin interpretation can be reasonably applied to the broad Balmer lines, suggesting that only a small amount of material should lie along the path of the photons. We can therefore conclude that a thorough investigation of the line intensity ratios along the profiles, which requires data with very high signal-to-noise ratios, will certainly provide further insight into the as yet not completely understood problem of the BLR structure.

4. Materials and Methods

The data and results presented in this paper are based on publicly available services and databases, such as the *VizieR* catalogue service¹ and the SDSS *SkyServer*². The plots have been produced with *TOPCAT*³ [26] and *XSPEC*⁴ software. Most of the illustrated measurements, involving line intensities and photometric colors, were directly extracted from public databases using their standard Web interfaces. Exceptions to the illustrated picture are the following:

- Measurements of emission-line normalized intensities were obtained from public SDSS spectra after extracting the BLR contribution by means of multi-Gaussian fits of the observed line profiles performed with *IRAF*⁵.
- Measurements of the ionized-plasma column density, obtained through XSPEC models combining a power-law emission, a soft X-ray thermal excess, an absorption contribution from neutral gas constrained by the H I column density within the Milky Way and an ionized absorber, applied to a 40 ks observation carried out by XMM Newton and reduced with the Science Analysis Software (SAS), version 10.
- 3. The models of SED in multi-frequency AGN data, obtained by applying Levemberg–Marquardt non-linear χ^2 minimization to combinations of cut-off power laws and black-body contributions.

Table 1 contains a summary of the data sources from which we obtained the SED points, while Table 2 reports a summary of the parameters used in the spectral energy distribution (SED) fitting models.

¹ http://vizier.u-strasbg.fr/viz-bin/VizieR

² http://skyserver.sdss.org/dr14/en/home.aspx ³ http://skyserver.sdss.org/dr14/en/home.aspx

³ http://www.star.bristol.ac.uk/mbt/topcat/

⁴ https://heasarc.gsfc.nasa.gov/xanadu/xspec/

⁵ http://iraf.noao.edu/

Instr. / Catalogue	$\log \nu$ (Hz)	Band	Reference	
NVSS	9.15	Radio	[27]	
IRAS	12.48-13.40	FIR	[28]	
WISE	13.13-13.94	FIR	[29]	
2MASS	14.14 - 14.38	NIR	[30]	
GALEX	15.00-15.30	UV	[31]	
XMM	16.86-17.68	X-ray	[32]	
INTEGRAL	18.70-19.00	Soft γ ray	[33]	
Fermi/LAT	23.00-26.00	γ ray	[34]	

Table 1. Data sources for the selection of multiple-wavelength spectral energy distribution (SED) points.

Table 2. Spectral energy distribution (SED) fitting models.

Object	Function	$\log v_{min}^{(a)}$	$\log v_{max}^{(a)}$	Norm.	Index ^(b)	Temp.	χ^2_{red}
3C 273	Power law	8.6	13.1	$2.30 imes 10^{-18}$	0.60	_	
	Power law	13.1	17.6	$1.31 imes 10^{-6}$	-0.31	-	
	Power law	17.1	22.6	$1.26 imes 10^{-23}$	0.70	-	
	Black body	-	-	$1.03 imes 10^{-12}$	-	25,800 K	1.11
NGC 1068	Power law	8.5	13.0	2.49×10^{-28}	1.52	-	
	Power law	12.8	17.5	7.068×10^2	-0.85	-	
	Power law	17.0	24.0	$4.82 imes 10^{-13}$	0.05	-	
	Black body	-	-	$1.04 imes 10^{-4}$	-	540 K	1.17

^(a) Logarithm of exponential cut-off frequency given in Hz; ^(b) The power-law index is given according to the notation $\nu F_V \propto \nu^a$.

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Author Contributions: G.L.M. is the contribution speaker, the main author of the text and the investigator of AGN multi-frequency properties; M.B. adapted the multi-frequency archive with timing capabilities; S.C. works on samples of NLS1 galaxies; A.C. contributed to the selection of all emission-line spectra with diagnostic lines from SDSS Data Release 14; S.C. supervised spectroscopic data reduction procedures; E.C. investigated advanced models of NLR emission lines; V.C. selected and analyzed the SDSS DR7 sample of NLS1; M.F. developed models of NLR structure; S.M. worked on multiple Gaussian fits and intensity ratios of optical lines; P.R. conceived the multi-frequency AGN archive and supervised the physical interpretation of optical spectra.

Conflicts of Interest: The authors declare no conflict of interest.

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Abbreviations

The following abbreviations are used in this manuscript:

- AGN Active galactic nucleus
- BLR Broad-line region
- FWHM Full width at half maximum
- NGC New General Catalogue
- NLR Narrow-line region
- PLTE Partial local thermodynamic equilibrium
- SED Spectral energy distribution

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Article Quasar Black Hole Mass Estimates from High-Ionization Lines: Breaking a Taboo?

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Abstract: Can high ionization lines such as CIV λ 1549 provide useful virial broadening estimators for computing the mass of the supermassive black holes that power the quasar phenomenon? The question has been dismissed by several workers as a rhetorical one because blue-shifted, non-virial emission associated with gas outflows is often prominent in CIV λ 1549 line profiles. In this contribution, we first summarize the evidence suggesting that the FWHM of low-ionization lines like H β and MgII λ 2800 provide reliable virial broadening estimators over a broad range of luminosity. We confirm that the line widths of CIV λ 1549 is not immediately offering a virial broadening estimator equivalent to the width of low-ionization lines. However, capitalizing on the results of Coatman et al. (2016) and Sulentic et al. (2017), we suggest a correction to FWHM CIV λ 1549 for Eddington ratio and luminosity effects that, however, remains cumbersome to apply in practice. Intermediate ionization lines (IP ~ 20–30 eV; AIIII λ 1860 and SiIII] λ 1892) may provide a better virial broadening estimator for high redshift quasars, but larger samples are needed to assess their reliability. Ultimately, they may be associated with the broad-line region radius estimated from the photoionization method introduced by Negrete et al. (2013) to obtain black hole mass estimates independent from scaling laws.

Keywords: ionization processes; emission line formation; atomic spectroscopy; supermassive black holes; emission line profiles; quasars

1. Introduction: Statement of the Problem

A defining property of type-1 quasars is the presence of broad and narrow optical and UV lines emitted by ionic species over a wide range of ionization potentials (IPs, [1]): high-ionization lines (HILs) involving IP > 50 eV, and low-ionization lines (LILs) from ionic species with IP < 20 eV. Over the years, it has turned expedient to consider the UV resonance line CIV λ 1549 as a representative of broad HILs. Lines do not all show the same profiles, and redshifts measured on different lines often show significant differences. Internal line shifts involving both broad lines in the optical and UV spectra of quasars have offered a powerful diagnostic tool of the quasar innermost structure since a few years after the discovery of quasars [2].

The CIV λ 1549 is a resonant doublet $({}^{2}P_{\frac{3}{2},\frac{1}{2}} \rightarrow {}^{2}S_{\frac{1}{2}})$ emitted by gas which is, at least in part, flowing out from a region within ~1000 gravitational radii from the central black hole (e.g., [3], for a review).

The occurrence of CIV λ 1549 large shifts constrains the suitability of the CIV λ 1549 profile broadening as a virial black hole mass (M_{BH}) estimator (see, e.g., [4,5], for reviews). Results at low-redshift obtained in the mid-2010s suggest that the CIV λ 1549 line is unsuitable for, at least, part of the quasar Population A sources (Section 4 [6]). A similar conclusion was reached at $z \approx 2$ on a sample of 15 high-luminosity quasars [7]. More recent work tends to confirm that the CIV λ 1549 line width is not straightforwardly related to virial broadening (e.g., [8]). However, the CIV λ 1549 line is strong and observable up to $z \approx 6$ with optical spectrometers. It is so highly desirable to have a consistent virial broadening estimate up to the highest redshifts that various attempts (e.g., [9]) have been done at "rehabilitating" CIV λ 1549 line width estimators to bring them in agreement with the width of LILs such as H β and MgII λ 2800 [10].

In this contribution, we briefly stress the importance of black hole mass estimates (Section 2) and recapitulate the basic method of $M_{\rm BH}$ estimates applied to large samples of quasars under the virial assumption (Section 3). To improve estimates that have been known to be plagued by systematic and statistical uncertainties as large as a factor ~100, we adopt the "rehabilitating" power of the quasar eigenvector 1 (E1; Section 4). Our analysis is then focused on virial broadening estimators (VBEs; Section 5) from: (a) LILs (IP < 15 eV: H β , MgII 2800); (b) HILs (IP > 40 eV: CIV λ 1549); (c) intermediate-ionization lines (IILs: SiIII] λ 1892, Al III λ 1860), for which we provide a brief summary of preliminary results. Reported results from our group were published during the last decade [11–14]. We finally suggest that "photoioionization" computations of $M_{\rm BH}$ (Section 6) may offer a solution to some of the problems associated with the use of an average scaling law.

2. Importance of Black Hole Mass Determination

The relevance of supermassive black hole M_{BH} estimates is not limited to the sake of a better understanding of quasars interpreted as accreting system. Black hole masses are a key parameter in the evolution of the galaxies and in cosmology as well. Massive, fast outflows are affected by the ratio of radiation to gravitational forces. They provide feedback effects to the host galaxy, and are even invoked to account for the M_{BH} -bulge velocity dispersion correlation [15–17]. The ratio between radiation and gravitation forces also influences broad-line region dynamics; lower column density material may flow out of the emitting region [14,18,19]. Black hole masses of high redshift quasars provide constraints on primordial black hole collapse (and references therein [20,21]). Overestimates by a factor as large as ~100 for supermassive black holes at high *z* may even pose a spurious challenge to concordance cosmology. Present-day estimates constrain the relation between the formation of the seed black holes and the collapse of the protogalaxy. Collapse of dark matter clumps yielding massive seeds that then rapidly grows by super-Eddington accretion appears necessary to explain the occurrence of supermassive black holes at very high redshifts.

3. Virial Black Hole Mass Estimates

The virial expression for the mass can be written as

$$M_{\rm BH} = f r_{\rm BLR} \delta v^2 / G,\tag{1}$$

where *f* is a factor of order unity reflecting geometry and dynamics of the broad line emitting regions, and r_{BLR} is a representative distance of the broad-line emitting region (BLR).¹ The virial broadening δv is provided by a measurement of the line profile width, which can be velocity dispersion σ , FWHM, or FWZI. The FWHM is by far the most handy measure employed as a VBE although some authors claim that a better estimator may be offered by the velocity dispersion [22]. The latter measure is, however, not defined in the case of Lorentzian profiles and not of easy interpretation in the case of

 $^{^{1}}$ The following analysis pertains only type 1 quasars showing broad (FWHM > 1000 km s⁻¹) emission in permitted lines.

shifted profiles. The virial assumption had a spectacular confirmation (albeit limited to few cases) by type-1 sources for which several lines were monitored: the response time of the BLR line and the line width were found to be correlated exactly as expected for Keplerian motion around a massive central object, with $\delta v \propto r^{-\frac{1}{2}}$ [23].

We can subdivide the estimates of the radius of the BLR (r_{BLR}) as primary and secondary. Primary determinations are obtained as the peak or centroid of the cross-correlation function between continuum and line light curves. Secondary determinations stem from the correlation between r_{BLR} and luminosity that has been derived from reverberation-mapped sources (e.g., [24–27]): $r_{BLR} \propto L^a$, $a \approx 0.5$ –0.65 [24,28]. The relation takes different form for different lines, and has been defined for H β and CIV. It then becomes possible to derive scaling laws for the black hole mass: $M_{BH} = M_{BH}(L, FWHM) = kL^a FWHM^b$. If a = 0.5, b = 2, the relation is consistent with the virial assumption (e.g., [29,30]), and the r_{BLR} scaling laws. If $a \neq 0.5$, $b \neq 2$ (e.g., [31–33]), the mass scaling law still provides M_{BH} estimates, but the accuracy of these estimates is likely to be sample-dependent (see Section 5.3).

The $M_{\rm BH}$ scaling laws provide a simple recipe usable with single-epoch spectra of large sample of quasars. Estimates of the Eddington ratio are derived by applying a bolometric correction to the observed luminosity. The correction is typically assumed a factor 10–13 from the flux λf_{λ} at 5100 Å (measured in erg s⁻¹) and 3.4–5 from λf_{λ} measured at 1450 Å in the UV, [34,35]). For the sake of the present review, we will stay with these constant corrections without forgetting that different bolometric corrections should be defined along the quasar "main sequence" (introduced in Section 4), and that the correction is most likely luminosity as well as orientation dependent [36].

Caveats

The estimate of the $M_{\rm BH}$ is based on several assumptions underlying reverberation mapping studies: among them, a compact region wetting the quasar continuum, and a fairly monotonic response of the line emitting gas. This latter assumption has been apparently challenged by the unpredicted behavior of NGC 5548 in 2014, where a time delay τ much shorter than expected was found when the source was in a high-luminosity state [37–39]. The physical origin of this behavior is not yet clear: in principle, shielding, optically thin gas, changing size of the continuum source could also give rise to a shorter (or a lack of) response to continuum change in the emitting line regions. In addition, periodic signals (in photometric and spectroscopic measurements) have been detected in a number of sources [40], including NGC 5548 (e.g., [41–45]). The origin of the periodicity is as yet unclear as well. A second, supermassive binary black hole has been suggested in some cases [41,46] but, in principle, black holes of masses much smaller than the primary could produce periodic photometric properties without leaving a detectable trace of their gravitational pull on the line profiles. As a matter of fact, the $r_{\rm BLR}$ -L scaling relation has a non-negligible intrinsic dispersion, which may be, at least in part, accounted for by a dependence of $r_{\rm BLR}$ estimates on the dimensionless accretion rate found in recent work [27].

The basic assumption underlying the search of virial broadening estimators is that a line (or line component) is symmetric and unshifted with respect to the quasar rest frame. Most line profiles are asymmetric, and shifted, but this has been ignored until recent years. If CIV FWHM is employed as a virial broadening estimator, the loss of information is so severe that the $M_{\rm BH}$ distribution as a function of redshift cannot be distinguished if masses obtained from random values of the FWHM (e.g., [47]) are used in place of the actual estimates.

A single value of the structure factor is obtained by scaling the M_{BH} to agree with the dynamical masses [48–52]. We expect that the geometry and dynamics of the BLR is related to the accretion mode, although it is as yet not unclear in which way. The structure factor is therefore likely to be different for different type-1 quasar populations. For Populations A and B (defined in Section 4), Collin et al. [53] find f(FWHM) \approx 2 and 0.5, respectively.
4. The Rehabilitating Power of Eigenvector 1

The eigenvector 1 (E1) was originally defined by a principal component analysis of ≈ 80 Palomar–Green quasars, and is associated with an anti-correlation between the strength of the FeII λ 4570 blend, measured by the flux ratio $R_{\text{FeII}} = F(\text{FeII}\lambda$ 4570)/ $F(\text{H}\beta)$ and the FWHM of H β . The E1 is (at the very least) an useful tool to organize quasar diversity through a sequence (the main sequence, MS) in the so-called optical plane of the E1, defined by FWHM(H β) vs. R_{FeII} . The 4D E1 parameters space includes two more parameters: the soft X-ray photon index Γ_{soft} associated with the accretion status, and the centroid at half maximum of CIV λ 1549 c(1/2) CIV measuring the amplitude of the high ionization outflow originating from the BLR [54]. More parameters, however, correlate with E1 (see Table 1 of Sulentic et al. [55] and Fraix-Burnet et al. [56]). Since the analysis of Boroson and Green [57], MS trends have been found in works dealing with a number of phenomenological and physical correlates [58–65]. Recently, large Sloan Digital Sky Survey (SDSS) data samples were involved [9,66–69]. The E1 "rehabilitating" power stems, in this context, by the ability to identify systematic trends that may be missed and confused as statistical errors if structurally different sources are dumped together in a single sample.

The trends along the MS allow for the definition of spectral types [70] in bins of $\Delta R_{\text{FeII}} = 0.5$ (A1, A2, A3, A4 from $R_{\text{FeII}} = 0$ to $R_{\text{FeII}} = 2$, tracing a change of L/L_{Edd} convolved with orientation, [71]) and FWHM (A for FWHM H $\beta < 4000 \text{ km s}^{-1}$, B1, B1⁺, B1⁺⁺ for broader sources in steps of 4000 km s⁻¹, tracing mainly a change in orientation). Quasars can be classified as belonging to two quasar Populations, A and B [72]. Population A (FWHM H $\beta < 4000 \text{ km/s}$) includes NLSy1s. Population A and B(roader) sources are most likely associated with a different accretion mode, since Population A are at $L/L_{\text{Edd}} > 0.2$ –0.3 which is the theoretical limit above and is an optically and geometrically thick advection-dominated accretion disk forms [73]. Population B of low L/L_{Edd} may be consistent with a flat α disk. The distinction between Populations A and B separates sources that are also called wind- and disk-dominated by Richards et al. [66], or Populations 1 and 2 by Collin et al. [53]. It supersedes the distinction between NLSy1s and rest of type-1 AGNs, which is based on a FWHM limit that has played an important historical role but is physically arbitrary. At low z (<0.7), Population A show low M_{BH} , high L/L_{Edd} Population B, high M_{BH} , low L/L_{Edd} , a reflection of the "downsizing" of nuclear activity: practically no black hole with very large mass ($M_{\text{BH}} \sim 10^9 - 10^{10} \text{ M}_{\odot}$) is accreting super-Eddington in the local Universe.

5. Virial Broadening Estimators along the Quasar MS

5.1. LILs: $H\beta$ and $MgII\lambda 2800$

The profiles of LILs like H β and MgII λ 2800 change along the quasar MS. If we want to extract a VBE, we have to consider the behavior of H β and MgII λ 2800 from extreme Population B to extreme Population A.

In Population B, the broad profiles of H β and MgII λ 2800 are most frequently redward asymmetric. Composite spectra of individual spectral types can be modeled by a broad Gaussian component (the BC, symmetric and unshifted, assumed to be the virialized component), and a redshifted very-broad Gaussian component. This latter component presumably associated with "perturbed" virialized motions or gravitational redshift in the inner BLR [74] systematically increases the line width with respect to the BC that provides the VBE.

At the other end of the MS extreme, Population A sources show narrower, Lorentzian-like profiles, slightly blueward asymmetric. The H β and MgII profiles can be modeled by Lorentzian functions plus a blueshifted excess modeled with a skewed Gaussian [12,13]. Both the LIL resonance line MgII λ 2800 and H β are affected by low-ionization outflows detected in the extreme Population A corresponding to spectral types A3–A4 [13]. However, in most of Population A, the LILs are dominated by a symmetric, virialized broad component [8,13,30,75].

We can define a parameter ξ yielding a correction to the observed profile to obtain a VBE, as follows:

$$\xi = \frac{FWHM_{\text{VBE}}}{FWHM_{\text{obs}}} \approx \frac{FWHM_{\text{BC}}}{FWHM_{\text{obs}}} \approx \frac{FWHM_{\text{symm}}}{FWHM_{\text{obs}}},$$
(2)

where the VBE FWHM can be considered best estimated by the FWHM of the broad component FWHM_{BC}, whose proxy can be obtained by symmetrizing the observed FWHM by various corrections described below. Even if the non-virial broadening mechanism is different along the E1 sequence, $0.75 \leq \xi \leq 1.0$ for both H β and MgII. This implies that, even if LILs are not always asymmetric, a modest correction is, on average, sufficient to retrieve a VBE from the observed FWHM. Table 1 reports the current estimates along with bibliographic references.

Table 1. The LIL and IIL ξ (Equation 2) factor for the spectral types of the quasar MS, listed for the spectral types (SpT) of highest occupation.

SpT	Нβ	MgII	AlIII	Ref.
A3-A4	0.8/0.9	0.75/0.8	1.0	[12,76]
A1-A2	1.0	1.0	1.0	[12,76]
B1	0.8	0.9	1.35	[13,45]
B1+/B1++	0.8	0.9	1.35	[13,45]

5.2. LIL VBE at High L

At low *z*, the quasar population reaches bolometric luminosity ~10⁴⁷ erg s⁻¹ at most. It is interesting to consider the LIL VBE behavior if we add to a local sample sources more luminous than this limit. Composite spectra are helpful to outline the LIL H β behavior over a wide luminosity range. Marziani et al. [11] computed composites in step of 1 dex for 6 dex in luminosity, joining an HE/ISAAC high-*L* sample (52 sources) and an SDSS sample from Zamfir et al. [77]. The H β line becomes broader with increasing *L* (over 43 < log *L* < 48.5 [erg/s]), but shapes are similar: the Population A/Population B differences are preserved at high *L* [11]. At the same time, the minimum FWHM(H β) increases with luminosity. This result is consistent with the virial assumption and with the *r*_{BLR} scaling law:

$$M_{\rm BH} \propto r_{\rm BLR} FWHM^2 \propto L^a FWHM^2 \propto (L/M)^{-1} L^{(1-2a)/2}.$$
(3)

The minimum, luminosity-dependent FWHM is obtained for a limiting Eddington ratio \approx 1. As a consequence, the Population A limit is also luminosity dependent.

Extracting a VBE from LIL H β at high-*L* is possible by employing different symmetrization methods:

- substitution of the BC in place of the full Hβ profile. This method requires a multicomponent maximum likelihood fitting, and is fairly reliable for Population A (even for extreme Population A) and Population B spectral types where the VBC is creating a profile inflection;
- symmetrization of the profile: FWHM_{symm} = FWHM-2 c(1/2);
- correction based on spectral type. In practice, this means to correct Hβ for Population B sources by a factor ξ as reported in Table 1.

All symmetrization methods were found to be equivalent at low- and high-L [45]. In other words, the H β profile shapes at high L are consistent with those at low-z, lower L (with some caveats, [11]).

5.3. CIV

Virial broadening estimators extracted from the HIL CIV λ 1549 along the E1 sequence are unfortunately not easy to define. If we scale the H β emission and overlay it on CIV λ 1549 (Figure 1), we obtain a strong excess of blueshifted emission. The profile can be interpreted as an almost symmetric,

unshifted "virialized" emitting region + an outflow/wind component that dominates in A3/A4 spectral types (e.g., [78–81]) and at high luminosity. The largest shifts of CIV λ 1549 centroid at 1/2 along the MS are found in Population A [4,6,14].



Figure 1. Left: the CIV profile of HE0512, a Population A quasar in the sample of Sulentic et al. [14]. The profile has been decomposed into a symmetric, unshifted component (the virialized component, thick black line) and a blue shifted component modeled by a skewed Gaussian (thick blue line). The flat-topped emission on the red side of CIV is mainly HeII λ 1640; **Right**: the 1900 blend for HE0512. The AlIII and SiIII] lines at 1860 Å and 1892 Å are unshifted and symmetric, and their FWHMs are in agreement with the H β and of the CIV virialized component profile. The thick black components can be thought as scaled H β profiles, as in the case of CIV. Note that AlIII λ 1860 lacks the blue-shifted excess of CIV. The thin line shows the CIII] λ 1909 profile whose intensity remains highly uncertain because of the heavy blending with FeIII emission.

The Vestergaard and Peterson [29] scaling laws

$$\log M_{\rm BH} = c + a \log L + b \log \rm FWHM, \tag{4}$$

assumed that the width of CIV and H β are equivalent, with $a \approx 0.5$, b = 2, and c an average constant difference between the 5100 and the 1450 luminosity. Considering the FWHM CIV as a proxy of FWHM H β causes a bias along the E1 sequence. Especially for extreme Population A, errors can be as large as 2 dex. Figure 2 is the statement of the ensuing BH mass taboo for CIV.

Figure 2 (based on the data of Sulentic et al. [6]) emphasizes a systematic trend for which a corrective could be estimated. To this aim, we defined a sample of RQ quasars with CIV λ 1549 observations including 70 sources at z < 1 [6], and 25 high-luminosity sources at z > 1.4 [14]. Matching H β data are available from our previous observations [11,82] or from published spectra. At $L > 10^{47}$ erg s⁻¹, high amplitude CIV 1549 blueshifts in both Population A and B are observed with median ≈ 3000 km s⁻¹ for Population A; two extreme cases involve CIV c(1/2) blueshift amplitude larger than 5000 km s⁻¹. Widespread powerful outflows are affecting both Population A and B sources [14,83] with worrisome implications for $M_{\rm BH}$ estimates from CIV λ 1549 FWHM.

Population A sources are more frequently selected at high *z*, high *L*. Figure 2 of [84] depicts what may be called an Eddington ratio bias: for a fixed mass, higher Eddington ratio sources are preferentially better sampled. In this way, the FWHM CIV (if left uncorrected) may lead to systematic over-estimates of M_{BH} by even one/two orders of magnitude, with the potential of creating the (spurious) result of a population of extremely massive black holes at high redshift (z > 2).



Figure 2. Bias of the M_{BH} estimates in the optical E1 plane FWHM(H β) vs. R_{FeII} , for the full sample of Sulentic et al. [6]. Grey symbols represent differences $|\Delta \log M| = |\log M(CIV) - \log M(H\beta)| \le 0.3$, blue dots $\Delta \log M > 0.3$, red circles $\Delta \log M < -0.3$. The size of the circles is proportional to the amplitude of the difference, with the largest blue circles showing CIV masses in excess by a factor 100. Mid-sized circles indicate a disagreement of a factor ≈ 10 . The labels identify the spectral types as described in Section 4 and the thick dot-dashed line at 4000 km s⁻¹ shows the boundary between Population A and Population B.

5.4. A CIVλ1549 VBE

The centroid shift at half-maximum c(1/2) is correlated with the FWHM in the CIV line, implying that the FWHM excess with respect to H β is associated with a blueshifted component, as shown in Figure 1.² The comparison of Figure 1 has been possible because we have H β observations for this source. However, H β observations are available only for a few hundred high-redshift quasars since the line is shifted in the IR.

Nonetheless, there have been several attempts in the last few years to define scaling laws that corrected for the non/virial contamination of CIV emission. A scaling law that assumes $M_{\rm BH} \propto$ FWHM^{0.5} [32] accounts for the over-broadening of Population A sources, but overcorrects for Population B [45]. In general, corrections dependent on $L/L_{\rm Edd}$ (which is correlated with FWHM of CIV, [85]) or an $L/L_{\rm Edd}$ proxy such as the SiIV + OIV]1400 blend/CIV 1549 ratio are promising [9] but tend not to work well for Population B. Empirical corrections based on CIV blueshift work fairly well for high-*L* sources only [83,86].

It is important to consider that there is a threshold in CIV shift amplitude (c(1/2)) at $L/L_{Edd} \approx 0.2$ [14]. There is a strong correlation with L/L_{Edd} if blueshifts are significant. There is also a weak but significant correlation with luminosity in the sample of Sulentic et al. [14]: the partial

² This happens also for MgII, but with much lower blueward displacements (few hundreds vs. few thousands km s⁻¹).

correlation coefficient between c(1/2) and $L(L/L_{Edd}$ hidden) is significant at about a 2σ confidence level. A multivariate analysis confirms the blueshift dependence on both L and L/L_{Edd} in that sample. The blueshift dependences are consistent with a radiation-driven outflow, with a slope ≈ 0.15 for log L, and slope ≈ 0.5 for L/L_{Edd} . A pure dependence on L arises for L/L_{Edd} in a small range. A strong dependence on L/L_{Edd} and a weak dependence on L can be obtained under a variety of physical scenarios. We expect that strength and form of L and L/L_{Edd} correlations are sample dependent [14].

Considering the threshold and the dependence on both *L* and L/L_{Edd} , using c(1/2) as a proxy for L/L_{Edd} , HIL CIV corrections based on c(1/2) and *L* reduce scatter to 0.33 dex with respect to H β estimates, providing an unbiased M_{BH} estimator. Unfortunately, the correction coefficients are different for Pops. A and B. For Population B, the correction is highly uncertain in the sample of Sulentic et al. [14], and a larger sample of sources is needed. In addition, (1) corrections derived by Coatman et al. [83] may be sample dependent because of the Eddington ratio bias; and (2) any correction based on c(1/2) requires a precise estimate of the quasar rest frame of reference, which is not easily obtained from UV data. A theoretical correction requires that c(1/2) CIV and ionization conditions in the BLR are accounted for, and has not been competed as yet. It is therefore not obvious whether M_{BH} estimates based on CIV can be reliable for large samples of quasars.

5.5. Intermediate Ionization Lines (IILs)

An alternative to the use of CIV is to resort to the emission lines of the 1900 Å blend, whose constituents are mainly the resonant doublet of AlIII at $\lambda 1860$ Å, and the intercombination lines SiIII] and CIII] at $\lambda 1892$ and $\lambda 1909$ Å, respectively (Table 1 of Negrete et al. [87] provides detailed information of the atomic transitions involved). We base our analysis on ≈ 80 objects of the CIV sample described in Section 5.3 for which the 1900 blend has been covered. CIII] $\lambda 1909$ measurements have serious problems: in Population A, CIII] is faint, and blended with strong and diffuse FeIII emission. In Population B, the CIII] $\lambda 1909$ line is affected by VBC. However, considering AlIII and SiIII] it is possible to obtain a VBE consistent with H β . To this aim is necessary to assume that FWHM AlIII 1860 = FWHM SiIII] 1892 i.e., to anchor the FWHM AlIII to FWHM SiIII]. A source of concern is that AlIII is a resonance doublet $({}^2P_{\frac{3}{2},\frac{1}{2}} \rightarrow {}^2S_{\frac{1}{2}})$ and part of its emission may originate in an outflow, where the AlIII, as does CIV, acts as a resonant scatterer of continuum photons. However, measured AlIII shifts are < 0.2 CIV shifts and the AlIII and SiIII] profiles looks fairly symmetric in the wide majority of cases (Figure 1 shows a typical case).

If we compare the FWHM of the IILs to H β , we find that it is in very good agreement with Population A. Population B IILs are narrower than H β , but a modest correction is needed to enforce an unbiased agreement: FWHM H $\beta_{BC} = 1.35$ FWHM AlIII_{*BC*}. In other words, $\xi \approx 1.35$ (Table 1) is needed to rescale the line widths to the one of H β BC assumed as a reference VBE.

6. Photoionization Masses

The ionization parameter U can be written as

$$U = \frac{Q(H)}{4\pi r_{BLR}^2 cn'},\tag{5}$$

where Q(H) is the number of ionizing photons, *n* the hydrogen density, and *c* the speed of light. The expression for *U* can be inverted to obtain the BLR radius

$$r_{\rm BLR} = \left(\frac{Q(H)}{4\pi U cn}\right)^{1/2}.$$
(6)

The radius r_{BLR} estimates from photo-ionization agree with $c\tau$ from reverberation mapping for a sample of 12 sources [76]. The photon flux $n \cdot U$ is estimated using diagnostic ratios involving AlIII 1860, SiIII] 1892, SiII 1816, CIV 1549, SiIV + OIV] 1400. The photoionization method provides an

unbiased estimator of r_{BLR} in the sample of Negrete et al. [76], but M_{BH} estimates at high L remain largely untested [88]. The photoionization method, in principle, could avoid the dispersion intrinsic to the scaling laws with L. To obtain accurate individual estimates of M_{BH} is then necessary to have an accurate knowledge of f and of orientation effects that influence the FWHM. Both are still poorly known at the time of writing.

7. Conclusions

Retrieving a VBE that is representative of the broadening due virial motions in the low-ionization part of the BLR is a major goal in order to reduce systematic and statistical effects in single-epoch $M_{\rm BH}$ determinations applied to large samples of quasars. The following remarks emerge from the present review.

- Low-ionization lines (Hβ, MgII 2800) provide reliable virial broadening estimators by applying corrections to the observed line width. The corrections depend on the spectral type along the E1 MS, but they are relatively small (less than 30%), and work up to the highest *L* of quasars.
- The HIL CIVλ1549 is not immediately providing a reliable virial broadening estimator. The profile
 is broadened by an excess emission on its blue side. The shift amplitude depends on both
 L/L_{Edd} and L. Large shifts are observed in Population A, with Eddington ratio above a critical
 L/L_{Edd} ≈ 0.2.
- It is possible to apply corrections to the observed CIV broadening, but they remain cumbersome even for Population A. Population B sources at low Eddington ratio require a different correction (still ill-defined by the analysis of Marziani et al. [45] as Population B sources are most affected by the Eddington ratio bias mentioned in Section 5.3).
- Preliminary results on the 1900 blend indicate that the IILs lines could provide a better choice than CIV; IIL FWHM measurements appear intrinsically more robust than those of CIV since they do not require corrections based on shift measurement with respect to rest frame. However, more data are needed to assess their reliability.

The ultimate solution may be to abandon scaling laws altogether and to attempt M_{BH} estimates on an individual basis, considering r_{BLR} from photoionization and $f = f(L/L_{Edd}, L, ...)$ (where ... indicates any relevant physical parameter, for example the spin parameter of the black hole) as well as orientation effects on the VBE along the E1 sequence.

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Abbreviations

The following abbreviations are used in this manuscript:

- BC Broad component
- BLR Broad line region
- E1 Eigenvector 1
- HE Hamburg-ESO
- HIL High ionization line
- IIL Intermediate ionization line

- IP Ionization potential
- ISAAC Infrared Spectrometer and Array Camera
- LIL Low ionization line
- FWHM Full width half maximum
- FWHM Full width zero intensity
- MS Main sequence
- NLSy1 Narrow-line Seyfert 1
- SDSS Sloan digital sky survey
- VBE Virial broadening estimator

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Project Report Photometric and Polarimetric Interpretation of Blazar AO 0235+164 Behaviour

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Abstract: Among the great number of controversial issues, the most topical one both for theoretical and observational astrophysics presently is a problem of active galactic nuclei investigation. To explain the behaviour of blazar AO 0235+164, which has been under observation at the LX200 telescope (SPbSU) since 2002, the method of analyzing developed by V.A. Hagen-Thorn and S.G. Marchenko was applied. It is based on the assumption that in the case of observational data lying on the straight line in the absolute Stokes parameters space {*I*, *Q*, *U*} (for polarimetry) and the fluxes space {*F*₁, ..., *F*_n} (for photometry), relative Stokes parameters and relative flux ratios stay unchanged , and consequently, only one source is corresponding to the variability of general value of flux. In this paper, the photometric and polarimetric interpretation of blazar behaviour is presented. Furthermore, the flux and flux–flux diagrams are obtained for three periods of object monitoring: 2006–2007 and 2008–2009 (outbursts) and 2009–2016 (decline with 2015 outburst).

Keywords: active galaxies; BL Lacertae objects; AO 0235+164; photometry; polarimetry

1. Introduction

One of the leading problems of both theoretical and observational astrophysics today is the determination and investigation of the nature of extragalactic objects' activity (active galactic nuclei, AGN). Currently, particular attention is paid to BL Lac objects considered to be compact active nuclei with a relativistic plasma jet closely aligned to the line of sight that provides non-thermal emission [1]. This type of orientation is able to clarify extremal properties of the sources such as violent variability and high degree of polarization.

One of the most intensively studied BL Lac objects is AO 0235+164. It was classified as an AGN in 1975 [2]; in the same work, the variability in the optical and radio wavebands was detected. Furthermore, the object reveals different values of redshift: z = 0.940 for emission light and z = 0.851 and z = 0.524 for absorption components, and for this reason there has been the conjecture of a gravitational microlensing mechanism [3]. Concerning the modern studies, the optical and radio light curves were analyzed, and the periodicity of the outburst on a ~5.7 yr time scale was suggested [4]. Moreover, intraday flux variability was also obtained at radio [5] and optical [6] wavebands and at high energies [7], while only at cm-wavelength could it be explained with interstellar scintillation.

The attention of this paper is focused on the blazar AO 0235+164. The monitoring of the blazar has been carried out on the telescopes of Saint Petersburg State University 16-inch Schmidt–Cassegrain reflector LX200 (Peterhof, Saint Petersburg) and 27.6-inch Cassegrain reflector AZT-8 (Crimean Astrophysical Observatory) since 2002. The long observational data set will be given and analyzed; the source properties will be revealed as well.

2. Analysis Method

Modern research is concerned with the estimation of active galaxies—and BL Lacertae objects particularly—as complicated systems where the underlying or "host" galaxy, inner emission region (accretion disk, broad line region (BLR), central engine), different parts of jet, etc. make contributions to the total flux. It is also considered that the source responsible for variations is located in the area close to the central engine. As active galaxies cannot yet be resolved into separated components by means of direct observations, the source properties giving an idea of the nature of nuclei activity can be revealed solely with indirect methods.

One of the essential source characteristics is the spectral energy distribution (SED). It can be obtained by gaining multicolour data on photometric variability. In that case, a long data set needs to be collected to judge the phase of object activity. Moreover, the most recognizable features of BL Lac objects' emission are high polarization degree (up to 60–70% at outbursts) [8] and variability of both polarization degree and angle.

To analyze the variable source behaviour, the method worked out by V. Hagen-Thorn and S. Marchenko was chosen. At its foundation lies the assumption that variability is caused by only one source. In order for the variations to imply only the total flux value changing while the relative Stokes parameters (for polarimetry) and relative SED (for photometry) stay unchanging, then in the space of absolute Stokes parameters {I, Q, U} or in flux space { $F_1, ..., F_n$ }, the observational data should lie on a straight line (Figure 1). Moreover, the inclination angles are relevant to relative Stokes parameters and relative flux ratios in the chosen wavebands.



Figure 1. Polarization behaviour in space of absolute Stokes parameters (left) and photometry behaviour in three-dimensional flux space (right).

The method's application to the observational data requires the solution of the inverse task. As it was proven in [9], the opposite is also true: the linear relation of fluxes points at unchanging SED. Therefore, in the case of observational data lying on a straight line in the given space, only one source is corresponding to AGN variability. More details can be found at [9,10].

3. Observational Data

The observational data was obtained at the telescopes of Saint Petersburg State University stations by CCD cameras ST-7XME (produced by SBIG). The CCD camera properties are reported in the Table 1.

Photometric data was gained by multicolour observations with *B*, *V* Jonhson and R_c , I_c Cousins filters; calculated accuracy was $\approx 0.03^m$ [11]. Polarimetric data was provided with Savart plates rotated

through 45° with respect to each other. Observations were made at R_c waveband (AZT-8) and at the broad band with the effective wavelength close to R_c filter (LX200). The error value was less then 2%.

Table 1. Characteristics of the ST-7XME CCD camera.

Pixel number	765×510
Pixel size	$9 \times 9 \ \mu m$
Linear size	6.9 imes4.6~mm
Dark current	$1 e^{-}$ /pixel/sec at $0 °C$
RON	15 e ⁻

3.1. Photometry

Following [12] we could define three or more dimensional flux spaces, where flux–flux diagrams are projections to the relevant planes. Due to additivity, the flux can be presented as a sum of variable term F^{var} and constant term F^c . In the case of a single source assumption with unchanging SED, flux ratios can be given as:

$$F_i^{var} / F_i^{var} = c_{ij}, \tag{1}$$

where $c_{ij} = const$. Comparing different wavebands it can be revealed that:

$$F_{i} = c_{ij}F_{i} + (F_{i}^{c} - c_{ij}F_{i}^{c}),$$
(2)

where F_i and F_i are observed fluxes. It is easily seen that the multicolour observations represented in this paper with four different wavebands can be used to divide the host galaxy and the variable source fluxes. Therefore, computing the inclination angles for the flux–flux dependencies, the flux ratios could be found, and according to them the supposition of colour changes or colour permanency could be done that gives an opportunity to consider the source behaviour.

For the blazar AO 0235+164, photometric data covering the period from 2006 to 2016 in filters B, V, R_c , I_c was analyzed. The light curves are represented in Figure 2. In order to convert magnitudes to fluxes, the conversion "magnitudes to fluxes" for given filters and CCD camera are presented in the Table 2 [11].

Thereby, the flux diagrams were plotted (Figure 3). It could be clearly seen that throughout the chosen period, two outbursts (violent flux increase) took place—in 2006–2007 and 2008–2009, when the magnitude was up to 13^m (Figure 2), and also the period of decline—2009–2016, presumably, with the outburst in 2015. All of the data were divided to these three periods and flux–flux diagrams were studied as projections on planes in flux space {*F*_B, *F*_V, *F*_R, *F*_I}. The same periods were used for polarimetry interpretation. The data is presented in Figures 4–6.

Table 2. Calibration formulae for filters B, V, R_c, I_c.

Band	Logarithm of Flux $F(\nu)$ (mJy)
В	$lgF_B = 6.629 - 0.4 B$
V	$lgF_V = 6.561 - 0.4 V$
R_c	$lgF_R = 6.489 - 0.4 R_c$
I_c	$lgF_I = 6.406 - 0.4 I_c$



Figure 2. Light curves in filters *B*, *V*, *R*_{*c*}, *I*_{*c*} for 2006–2016.



Figure 3. Flux diagrams in filters B, V, R_c , I_c for 2006–2016.



Figure 4. Flux–flux diagrams for outburst 2006–2007.

• •

8 10 12

4 6

F. [mJv]



Figure 5. Flux–flux diagrams for outburst 2008–2009.



Figure 6. Flux-flux diagrams for decline 2009-2016.

3.2. Polarimetry

Polarimetric observations at the SPbSU station are dedicated to investigating the linear polarization— the Q and U Stokes parameters. The I parameter is considered to be equal to the F_R flux value. For data analysis, the same assumption and processing methods were used. As the Stokes parameters and photometric fluxes are additive, the observed values could be divided to the constant and variable parts: Q^c , U^c , and I^c , and Q^{var} , U^{var} and I^{var} . Then, the relative Stokes parameters p_x , p_y could be introduced:

$$p_x = p \cos 2\theta, p_y = p \sin 2\theta, \tag{3}$$

where *p* is a polarization degree: $p = (p_x^2 + p_y^2)^{1/2}$, θ is a polarization angle. p_x , p_y are considered to be constant. The observed Stokes parameters could be presented as:

$$\begin{cases} Q = p_x I + (Q^c - p_x I^c) \\ U = p_y I + (U^c - p_y I^c) \end{cases}$$
(4)

The system apparently describes the equation of a straight line, where p_x , p_y are direction tangents. To investigate the polarimetric behaviour in the case of the chosen method, the projections Q(I) and U(I) were plotted and studied for each period. The data is shown in Figures 7–10.



Figure 7. Absolute Stokes parameters behaviour (left) and Q(I) and U(I) (right) at the first period.



Figure 8. Absolute Stokes parameters behaviour at different flashes (left) and Q(I) and U(I) (right) at the second period.

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Figure 9. Q(I) and U(I) (left) and Q(I) and U(I) for the flash (right) at the third period.



Figure 10. Absolute Stokes parameters behaviour at the third period (the flash is marked).

4. Results

Within the confines of the method, the obtained flux–flux curves could be approximated with a linear equation of type: $X = \alpha Y + \beta$, where *X* and *Y* are fluxes *F_i* and *F_j* or the Stokes parameters. As values of *X* and *Y* are loaded with errors, they are considered as random values distributed equally and not correlated. Thus, the method of orthogonal regressions of A. Wald described in [13] was applied. To calculate the inclination angle α , the next expression was used:

$$tg2\alpha = 2\frac{\sum_{k=1}^{n} (x_k - \bar{x})(y_k - \bar{y})}{\sum_{k=1}^{n} (x_k - \bar{x})^2 - \sum_{k=1}^{n} (y_k - \bar{y})^2},$$
(5)

where x_k and y_k are values of X and Y, respectively , and \overline{x} and \overline{y} are average values. Therefore, the inclination angles were calculated by photometric diagrams $F_B(F_I)$, $F_V(F_I)$, and $F_R(F_I)$ (see Table 3) and by polarimetric diagrams Q(I) and U(I) (see Table 4) for three periods of the blazar activity.

	$B(I_c) \\ \delta B(I_c)$	$\frac{V(I_c)}{\delta V(I_c)}$	$\frac{R(I_c)}{\delta R(I_c)}$
Outburst 2006–2007	$0.151 \\ \pm 0.034$	$0.280 \\ \pm 0.053$	0.455 ± 0.029
Outburst 2008–2009	0.162 ± 0.007	$0.308 \\ \pm 0.043$	0.485 ± 0.006
Decline 2009–2016	0.126 ± 0.021	0.287 ± 0.005	0.455 ± 0.053

Table 3. The inclination angles for photometric data.

Table 4. The inclination angles for polarimetric data.

	$Q(I) \\ \delta Q(I)$	$U(I) \\ \delta U(I)$
Outburst 2006–2007	-0.156 ± 0.013	$\begin{array}{c} -0.130 \\ \pm 0.026 \end{array}$
Outburst 2008–2009	-0.284 ± 0.011	-0.052 ± 0.011
Decline 2009–2016	0.092 ±0.014	-0.008 ± 0.014

5. Discussion

From the presented flux-flux diagrams and polarimetric dependencies it is obvious that Choloniewski's method, extended by Hagen-Thorn and described in Section 2, could be applied to the interpretation of the behaviour of blazar AO 0235+164. The possibility of the good photometric diagrams' approximation by straight lines reveals that within the two separated outbursts—2006–2007 and 2008–2009—fluxes ratios stayed unchanging, the object does not evince colour variability, and consequently, SED was permanent as well.

Furthermore, the comparison between the inclination angles at the different periods gives an approximate equality of all dependencies (the same filters are meaning) with the given accuracy. This could be mentioned as an indirect indication of the fact that physical properties of variable source were constant throughout the given event. In spite of possible multicomponent structure of radiating center, we could assume that the main contribution to the emission was made by a single source with a lifetime not less than the event duration. However, the data of decline period 2009–2016 noticeably differs from outbursts. It could be explained, firstly, by the lack of observations due to technical difficulties, and secondly, by gaining low fluxes with greater errors (and therefore less-accurate approximation). It could also be supposed that at the decline period the intensity of the variable source falls and the multicomponent structure displays changing SED.

According to the theory, the inclination angles of Q(I) and U(I) dependencies are equal to the relative Stokes parameters p_x , p_y . Due to the good agreement with the linear relations, it could be claimed that these parameters also remain unchanging for each period. Moreover, the average polarization degree p could be found for the different events: 21%, 29%, and 9%. The striking deviations presumably represent the occurrence of rapid polarization state variations that conform well to the theoretical models. It should be mentioned that extrinsic polarization makes a contribution to the total value as well. So, the polarization carried by ISM and AGN components (e.g., accretion disk or dust torus) should be also taken into account. The degree of constant polarization could be

calculated with system (4). According to the observation process, the value I^c is equal to F_R^c ; the last value could be found by the flux–flux diagrams. Therefore, the permanent contribution to the polarized flux does not exceed several percent. Additionally, the instrumental polarization is not greater than 2%. These values are negligible compared with variable polarization degree—up to 40% at outbursts. In summary, the interval of the variable source polarization degree changes is approximately 0–40%; the polarization angle also changes within all possible values, and it is essential to note that evident correlation has not yet been revealed.

Presently, the work is still in progress. To clarify the object behaviour, we are going to find the coefficients of correlation between blazar intensity and its polarization. The time lag between polarized and unpolarized fluxes observation would reveal the polarization origin—inner of extrinsic (e.g., the reflection on the dust torus)—and according to that the conclusions regarding polarization nature could be drawn.

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Review



Symmetric Atom–Atom and Ion–Atom Processes in Stellar Atmospheres

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Abstract: We present the results of the influence of two groups of collisional processes (atom–atom and ion–atom) on the optical and kinetic properties of weakly ionized stellar atmospheres layers. The first type includes radiative processes of the photodissociation/association and radiative charge exchange, the second one the chemi-ionisation/recombination processes with participation of only hydrogen and helium atoms and ions. The quantitative estimation of the rate coefficients of the mentioned processes were made. The effect of the radiative processes is estimated by comparing their intensities with those of the known concurrent processes in application to the solar photosphere and to the photospheres of DB white dwarfs. The investigated chemi-ionisation/recombination processes are considered from the viewpoint of their influence on the populations of the excited states of the hydrogen atom (the Sun and an M-type red dwarf) and helium atom (DB white dwarfs). The effect of these processes on the populations of the excited states of the hydrogen atom (atom and an M-type red dwarf) and helium atom (DB white dwarfs). The effect of these processes on the populations of the excited states of the hydrogen atom has been studied using the general stellar atmosphere code, which generates the model. The presented results demonstrate the undoubted influence of the considered radiative and chemi- ionisation/recombination processes on the optical properties and on the kinetics of the weakly ionized layers in stellar atmospheres.

Keywords: atomic processes; molecular processes; radiative transfer; absorption quasi-molecular bands; sun:atmosphere; sun:photosphere; stars:atmospheres; white dwarfs

1. Introduction

Atomic and molecular data play a key role in many areas of science like atomic and molecular physics, astrophysics, nuclear physics, industry, etc. [1-9]. The interpretation of interstellar line spectra with radiative transfer calculations usually requires spectroscopic data and collision data (e.g., atomic parameters, cross sections, etc.) [10,11]. Determination of accurate fundamental stellar parameters is one of the most important of today's tasks and this area of fundamental science is very important and still current. For example, the atomic and molecular data are important for development of atmosphere models of solar and near solar type stars and for radiative transport investigations as well as an understanding of the kinetics of stellar and other astrophysical plasmas [12]. Available LTE codes for stellar atmosphere modelling like ATLAS [13,14], MARCS [15] as well as NLTE codes like e.g., PHOENIX (see e.g., [16,17]) and TLUSTY [18,19] require the knowledge of atomic and molecular data. In addition, spectrum synthesis codes (e.g., SYNTHE, SYNSPEC) for radiative transfer and spectra depend on these as input parameters [20]. Such atomic data and processes are also important in modelling early Universe chemistry (see [21]). Evaluation of chemical abundances in the standard Big Bang model are calculated from a set of chemical reactions for the early universe and among them are very important reactions with species like H, H₂⁺ and also different Rydberg atoms [22]. The highly excited atomic states are named 'Rydberg states', and the atoms in such states are called 'Rydberg atoms'. Strictly speaking, only highly excited states, should be counted among the Rydberg

states. However, in practice, an atom $A^*(n, l)$ is being treated as a Rydberg atom if $(n - n_0) \ge 4$, where n_0 is the principal quantum number of the outer shell of the atom A in its ground state, and in the case of an atom $\text{He}^*(n)$ —for any $n \ge 3$. It should be noted that, at the present time, even in the laboratory experiments, Rydberg atoms with n close to 10^2 are being explored, while the astrophysicists observe the radiation of atoms from the states with n close to 10^3 . With a change of n, the parameters characterizing Rydberg states may change by orders of magnitude.

The content of this article is distributed in four sections. The first is devoted to the detailed description of the processes, and the corresponding methods of the determination of such processes coefficients and parameters. Sections 2 and 3 show the existing theoretical results concerning the investigated processes, their role in the low temperature layers of stellar atmospheres, as well as the methods of the investigation of such processes. Finally, at the end of this article, the current research and directions of further research are summarized.

In a series of papers [23–27], two groups of atom–Rydberg atom and ion–atom collisional processes have been studied from the point of view of their effect on the optical and kinetic properties of weakly ionized laboratory and astrophysical plasmas.

The first group of them includes chemi-ionization and chemi-recombination processes of the type

$$A^*(n) + A \iff e + A + A^+, \tag{1a}$$

$$A^*(n) + A \Longleftrightarrow e + A_2^+, \tag{1b}$$

where $A^*(n)$ is an atom in a highly excited (Rydberg) state with a principal quantum number $n \gg 1$, and e is a free electron. The processes caused by the action of the resonant energy exchange mechanism inside the electron component corresponding to the atomic–atomic or electron–ion–atomic system are considered [28]. These processes are illustrated in the Figure 1, where Figure 1a schematically shows the geometry of the collision $A^*(n) + A$, and Figure 1b—the essence of the resonant mechanism is that the transition of an external weakly bound electron of the system (e_n) with energy $\varepsilon_n < 0$ to a free state with energy $\varepsilon_k > 0$ is accompanied by a transition of the subsystem $A + A^+$ from the first excited molecular state with energy $U_2(R)$ to the ground state with energy $U_1(R)$.



Figure 1. (a) schematic illustration of $A^*(n, l) + A$ collision (1) within the domain of internuclear distances $R \ll r_n$, where r_n is the characteristic radius of Rydberg atom $A^*(n, l)$; (b) schematic illustration of the resonance mechanism.

The second group of processes includes radiation processes of the type:

$$\varepsilon_{\lambda} + A_2^+ \iff A + A^+$$
 photodissociation/association, (2a)

$$\varepsilon_{\lambda} + A^{+} + A \iff A + A^{+}$$
 radiation charge exchange, (2b)

where ε_{λ} stands for the energy of a photon with a wavelength λ , A and A^+ —an atom and its positive ion in the ground states, and A_2^+ —a molecular ion in the ground electronic state. These processes are illustrated in the Figure 2, where $U_1(R)$ and $U_2(R)$ represent the adiabatic potential curves of the ground and first excited electronic state of the ion, and R stands for the internuclear distance in atomic units. This figure shows that the radiative processes under study represent the result of transitions, with the emission or absorption of a photon, between the mentioned molecular electronic states.

The effect of chemi-ionization and chemi-recombination processes (1) can be estimated by comparing their intensities with the intensities of known concurrent ionization and recombination processes, namely:

$$A^+ + e \Longrightarrow \varepsilon_{\lambda} + A^*(n), \tag{3}$$

$$A^+ + e + e \iff A^*(n) + e. \tag{4}$$



Figure 2. The schematic presentation of the photo-dissociation/association processes Equation (2a) and free–free processes Equation (2b): *R* is the internuclear distance, $U_1(R)$ and $U_2(R)$ are the potential energy curves of the initial (lower) and final (upper) electronic state of molecular ion A_2^+ , and $h\nu$ is the photon energy.

The influence of radiation processes (2) can be analysed by comparing with the intensities of known concurrent radiative processes, namely:

$$A^{+} + e \Longleftrightarrow \varepsilon_{\lambda} + A^{+} + e, \tag{5a}$$

$$A^+ + e \Longleftrightarrow \varepsilon_{\lambda} + A^*, \tag{5b}$$

$$A + e \Longleftrightarrow \varepsilon_{\lambda} + A + e, \tag{6}$$

$$A + e \iff \varepsilon_{\lambda} + A^{-}, \tag{7}$$

where A^- is stable negative ion.

In connection with astrophysical plasmas, two cases are considered:

- the case of hydrogen, when A = H(1s) and $A^+ = H^+$,
- the case of helium, when $A = \text{He}(1s^2)$ and $A^+ = \text{He}^+(1s)$.

For the solar atmosphere, *A* usually denotes atom H(1s) and $A^+ = H^+$, and, for the case of helium-rich white dwarf atmospheres, *A* denotes He(1s²) and $A^+ = \text{He}^+(1s)$.

Chemi-ionization/recombination processes are very important for the evolution of Universe in the early epochs. The most important process is hydrogen recombination, and Rydberg states can play important roles in this process [22].

2. Chemi-Ionization and Chemi-Recombination Processes

The region of importance of chemi-ionization processes and production of Rydberg atoms i.e., chemi-recombination is in the cool dwarf stars and, especially, cool white dwarfs. Spectroscopic observations of cool white dwarfs [29] have demonstrated that white dwarfs with temperatures less than 6100 K are found to display significant flux deficits that are not predicted by the current WD model. One can suggest that such process may be absorption by atoms and molecules in highly excited Rydberg states.

Recent research of the atmospheres of cooling stars such as white dwarfs pointed out an anomaly in light emission of Rydberg atom with n = 10 and tabular lifetime $\tau \sim 10^{-6}$ s. The lines of the corresponding infra-red transitions have not been observed [30]. Let us note that it is just these states that correspond to the maximal values of chemi-ionization rate coefficients. According to the observational data, we have that, under such conditions $N_0 \ge 10^{17}$ cm⁻³ and $N^* \ge 10^{13}$ cm⁻³, where N_0 and N^* are the densities of the ground state and Rydberg atoms. It is not difficult to estimate that the probability of a Rydberg atom being extinguished through the chemi-ionization channel is comparable to the probability of its radiative decay.

Let us not forget the importance of chemi-ionization processes and production of Rydberg atoms in chemistry, physics and related branches of science [28,31,32].

In this article, the current state of research of the processes in atom–Rydberg atom collisions is presented. The principal assumptions of the model of such processes are based on the dipole resonance mechanism.

The chemi-ionization and inverse chemi-recombination processes (1) can be considered from the point of view of their effect on the population of the excited states of the hydrogen atoms in solar and cold-star atmospheres, as well as on the population of excited states of the helium atoms in DB white dwarfs. Comparative analysis of the influence of these and concurrent processes (3) and (4) can be presented by the values of the following parameters:

$$F_{phr}^{(ab)}(2,8) = \frac{\sum\limits_{n=2}^{8} I_r^{(ab)}(n,T)}{\sum\limits_{n=2}^{8} I_{phr}^{(ab)}(n,T)}, \quad F_{eei}^{(ab)}(2,8) = \frac{\sum\limits_{n=2}^{8} I_r^{(ab)}(n,T)}{\sum\limits_{n=2}^{8} I_r^{(eei)}(n,T)}, \quad F_i(n,T) = \frac{I_{ci}(n,T)}{I_{i;ea}(n,T),}$$
(8)

where $I_{ci}(n, T)$, $I_{i;ea}(n, T)$, $I_r^{(ab)}(n, T)$, $I_{phr}^{(ab)}(n, T)$ and $I_r^{(eei)}(n, T)$ are the fluxes caused by ionization and recombination processes (1), (3) and (4).

2.1. Solar Atmosphere

As a necessary step to improve the modeling of the solar photosphere, as well as to model atmospheres of other similar and cooler stars where the main constituent is also hydrogen, it is required to take into account the influence of all the relevant collisional processes on the excited-atom populations in weakly ionized hydrogen plasmas. This is important since a strong connection between the changes in atom excited-state populations and the electron density exists in weakly ionized plasmas.

It is a fact that, with an increase of the electron density, caused by a growth of the excited hydrogen atom population, the rate of thermalization by electron–atom collisions in the stellar atmosphere will become higher. A consequence will be that the radiative source function of the line center will be more closely coupled to the Planck function, making the synthesized spectral lines stronger for a given model structure, affecting the accuracy of plasma diagnostics and determination of the atmospheric pressure.

The theoretical investigation of the processes (1) started in [33] for the hydrogen symmetric case A = H. Although some of the chemi-ionization processes in atom–Rydberg atom collisions had already been described in [33], their intensive astrophysical research began somewhat later. From the astrophysical point of view, in Ref. [34], an investigation was started for the chemi-recombination processes of the photosphere and the lower chromosphere of the Sun, where $4 \le n \le 8$. Further research of Mihajlov and coworkers continued in [35] on the chemi-ionisation processes in $H^*(n \ge 2) + H(1s)$ collisions and inverse recombination in the photosphere and the lower chromosphere of the Sun.

The partial rate coefficients for the chemi-ionization processes (1a,b) are determined by expressions

$$K_{ci}^{(a,b)}(n,T) = \int_{E_{min}(n)}^{\infty} v\sigma_{ci}^{(a,b)}(n,E)f(v;T)dv,$$
(9)

where $\sigma_{ci}^{(a,b)}(n, E)$ is cross section, v is the atom–Rydberg–atom impact velocity, f(v; T) is the velocity distribution function for the given temperature T, and $E_{min}(n)$ is determined by the behavior of the potential curve $U_2(R)$ and the splitting term as presented in [35].

Under the conditions that exist in the solar atmosphere, the chemi-recombination rate coefficients can be presented over chemi-ionization rate coefficient

$$K_{ci}^{(a)}(n,T) \cdot N_n N_1 = K_{cr}^{(a)}(n,T) \cdot N_1 N_{ai} N_e,$$
(10)

where N_1 and N_n denote the densities of ground- and excited-state of atoms, respectively, while N_{ai} is the densities of ions.

Using partial rate coefficients $K_{ci,cr}^{(a,b)}(n,T)$, we can determine the total one,

$$K_{ci,cr}(n,T) = K_{ci,cr}^{(a)}(n,T) + K_{ci,cr}^{(b)}(n,T),$$
(11)

which characterizes the efficiency of the chemi-ionization/recombination processes (1a,b) together. The total chemi-ionization and chemi-recombination fluxes are $I_{ci}(n, T) = K_{ci}(n, T) \cdot N_n N_1$, $I_{cr}(n, T) = K_{cr}(n, T) \cdot N_1 N_i N_e$.

Figure 3a,b show the values of the total chemi-ionization and recombination rate coefficients Kci(n, T) and Kcr(n, T) in the region $2 \le n \le 8$ and $5000 \text{ K} \le T \le 10,000 \text{ K}$ obtained in [35]. From the figure, it follows that the maximum values of the hydrogen Kci(n, T) lies at n = 5 for all temperatures (except for T = 5000 K, where maximum value of Kci(n, T) are at n = 4), and maximum values for the chemi recombination rate coefficient Kcr(n, T) shifts to higher n (from 3 to 5) when temperature increases. The relative importance of a particular channel ('a' and 'b') for the chemi-ionization and chemi-recombination processes (1a,b) is shown in Figure 4. Branch coefficient $X(n;T) = K_{ci,cr}^{(b)} / Kci(n, T)$ present the ratio of rate coefficient for processes (1b) and total ((1a) + (1b)). From Figure 4, one can see that, for lower temperatures, i.e., 5000 K, processes (1b) are dominant while importance of processes (1a) increase with the increase of temperature as expected.



Figure 3. (a) total chemi-ionization rate coefficients Kci(n; T; H) with $5000 \text{ K} \le T \le 10,000 \text{ K}$ and for principal quantum numbers n = 2-10; (b) same as in (a) but for the inverse recombination coefficients Kr(n; T; H) (data taken from [35]).



Figure 4. Same as in Figure 3 but for the branch coefficient X(n;T;H), characterizing the relative importance of the particular channel ('a' and 'b') for the chemi-ionization and chemi-recombination processes (1a,b).

From the astrophysical point of view, the results have shown that, in some parts of the solar atmosphere, chemi-recombination processes (1) can dominate with respect to the photo-recombination process (3) and their intensity can be close to the intensity of triple electron-electron-ion recombination processes (4). The behavior of quantity $F_{phr}^{(ab)}$ as a function of *h* is shown in Figure 5a. The behavior of the quantity $F_{eei}^{(ab)}$ as a function of height *h* is shown in Figure 5b from [35]. One can see that the considered chemi-recombination processes dominate with respect to the concurrent electron-electron-ion recombination processes within the region $100 \,\mathrm{km} \le h \le 650 \,\mathrm{km}$ and significantly influence the optical properties of the solar photosphere. Namely, if we take the considered processes into account, we will improve the modeling of the solar photosphere, as the model atmospheres of other similar and cooler stars. If we do not take into account all relevant collisional processes on the excited-atom populations in weakly ionized plasmas, the corresponding electron density will be less accurate. For example, as stated in Reference [26], an increase of the electron density, caused by a growth of the excited hydrogen atom population, will result in a higher rate of thermalization by electron-atom collisions in the stellar atmosphere. As a consequence, the radiative source function of the line center will be more closely coupled to the Planck function, and the synthesized spectral lines will be stronger for a given model structure. This will influence on the accuracy of plasma diagnostics and determination of the atmospheric pressure.

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Domination of the chemi-recombination processes with $2 \le n \le 8$ over the electron–ion photo–recombination processes is confirmed in a significant part of the photosphere (-50 km $\le h \le 600$ km). Thus, it is proofed that these processes are important for non-LTE modeling of solar atmosphere



Figure 5. (a) parameter $F_{phr}^{(ab)}(n, T)$ (8) as functions of the height *h* of the solar atmosphere; (b) same as in (a) but for for the parameter $F_{oei}^{(ab)}(n, T)$ (8).

2.2. Atmospheres of the DB White Dwarfs.

The chemi-ionization and chemi-recombination processes and consequently Rydberg atoms are of importance for cool stars and, first of all, for cool white dwarfs. Recently, a new effect has been noticed from Spitzer observations of cool white dwarfs [29]. Namely, these observations have demonstrated that some white dwarfs with T < 6100 K are found to display significant flux deficits in Spitzer observations (see [29]). These mid-IR flux deficits are not predicted by the current white dwarf models including collision induced absorption due to molecular hydrogen. This fact implies that the source of this flux deficit is not standard molecular absorption but some other physical process. It is possible that such process may be absorption by atoms and molecules in highly excited Rydberg states.

In the helium case of chemi-ionization and chemi-recombination processes, the situation turns out to be similar to the hydrogen case. This conclusion is based on the results obtained by Mihajlov and coworkers in [36]. The influence of symmetrical chemi-ionization and chemi-recombination processes on the helium atom Rydberg states population in weakly ionized layers of helium-rich DB white dwarfs has been investigated in [36].

Figure 6a,b show the obtained values of the total chemi-ionization and recombination rate coefficients Kci(n, T) and Kcr(n, T) in the region $3 \le n \le 10$ and 10,000 K $\le T \le 30,000$ K for helium plasma. The dependence of the both coefficients Kci(n, T) and Kcr(n, T) on the quantum number decreases with the increase of the temperature. Regarding the importance of the particular channel ('a' and 'b') for the chemi-ionization and chemi-recombination processes (1a,b), as can be seen in Figure 6b, conclusions are the same as for the chemi-ionization rate coefficients.

From the astrophysical viewpoint, chemi-ionization and chemi-recombination contribution to the Rydberg state populations have been compared with electron–electron–ion recombination, electron–excited atom ionization, and electron–ion photorecombination processes for *n* from 3 to 10, in helium-rich DB white dwarf atmosphere layers with logarithm of gravity log g = 7 and 8 and effective temperature $T_{eff} \leq 20,000$ K.



Figure 6. (a) total chemi-ionization rate coefficients Kci(n; T; He) with 10,000 K $\leq T \leq$ 30,000 K and for principal quantum numbers n = 3-10; (b) same as in (a) but for the inverse recombination coefficients Kr(n; T; He) (data taken from [36]).

Some of the results obtained for the helium case [36] are illustrated by the Figure 7a,b, which are related to the recombination processes in the DB white dwarf atmosphere with an effective temperature $T_{eff} = 12,000$ K. The values of the parameters $F_{phr}^{(ab)}(n, T)$ and $F_{eei}^{(ab)}(n, T)$ are given as functions of the logarithm of the Rosseland optical depth.

The results from [36] undoubtfully show that, for the lower temperatures, the chemi-ionization/ recombination processes (1) are absolutely dominant over electron-excited atom ionization (3) and electron–electron–atom recombination (4) processes, for n = 3, 4, and 5 for almost all log $\tau < 0$ values. For n = 6, 7, and 8 and in the same log τ range, processes (1a,b) are comparable with processes (3) and (4). It is concluded that processes (1a,b) can be dominant ionization/recombination mechanisms in helium-rich DB white dwarf atmosphere layers for log g = 7 and 8 and $T_{eff} \le 20,000$ K and have to be implemented in relevant models of weakly ionized helium plasmas.



Figure 7. (a) parameter $F_{eei}^{(ab)}(n, T)$ (8) as a function of the logarithm of the Rosseland optical depth log τ ; (b) as in (a) but for the parameter $F_{phr}^{(ab)}(n, T)$ (8).

2.3. Chemi-Ionization Processes in Solar and DB White-Dwarf Atmospheres in the Presence of Mixing Channels

In the recent paper of Mihajlov et al. [37], two kinds of atomic collision processes have been considered that simultaneously occur in the stellar atmospheres and influence each other (see Figure 8). This is about the chemi-ionization processes (1) and (n-n')-mixing i.e., excitation-deexcitation processes

$$A^*(n,l) + A \Longrightarrow A + A^*(n',l), \quad (n-n')-\text{mixing}, \tag{12}$$

where *A* are atoms in their ground states, $A^*(n, l)$ is an atom in a highly excited (Rydberg) state with the principal quantum number $n \gg 1$ and orbital quantum number l and A = H or He.



Figure 8. The schematic presentation of the chemi-ionization and (n-n')-mixing processes.

It was expected that process (12) would reduce the impact of chemi-ionization processes (1), as can be seen from the data of Mihajlov et al. (see Figure 9). The arrows in Figure 9a,b illustratively present that reduction of rate coefficients.

Mihajlov and coworkers have shown that, for the lower temperatures, the chemi-ionization processes are still dominant over electron-excited atom ionization processes ($A^*(n) + e \rightarrow A^+ + e + e$), for n = 3, 4, and 6 almost in the whole observed atmosphere which is illustrated by Figure 10 from [37]. For n = 6, 7, and 8 in the whole observed atmosphere, chemi-ionization processes are comparable with electron-excited atom ionization processes. This is illustrated in Figure 10a, for WD with $T_{eff} = 12,000$ K (helium case) and in Figure 10b for solar photosphere (hydrogen case).

From the results [37,38], it follows that processes (1) are significant for such hydrogen and helium plasmas with the ratio $Ne/Na < 10^{-3}$, where Ne and Na are the free electron and ground state atom density. Accordingly, these processes are significant for the stellar atmospheres that contain the corresponding weakly ionized layers.



Figure 9. (a) comparison of the calculated values of rate coefficients of the chemi-ionization processes (1a,b) with and without inclusion of (n-n')-mixing process, case A = H; (b) same as in (a) but for case A = He.



Figure 10. (a) parameter $F_i(n, T; He^*(n))$ (ratio of fluxes generated in atom–Rydberg–atom and electron-excited atom impact ionization) as a function of the logarithm of Rosseland optical depth log(τ), for principal quantum numbers n = 3–10, with $T_{eff} = 12,000$ K and log g = 8: wide line—present calculation; tiny line—calculation from [35]; (b) parameter $F_i(n, T; H^*(n))$ as a function of the height h, for principal quantum numbers n = 3–8, for model of solar photosphere [39]: wide line—present calculation; tiny line—calculation from [36].

2.4. The Atmospheres of Late Type Dwarfs (M Red Dwarfs)

The general stellar atmosphere code PHOENIX [10] has the advantage that, apart from solving the atmospheric structure, it also calculates output spectra. A good example of testing and application of PHOENIX code are processes (1) that influence the excited state populations and the free electron density, and also influence the atomic spectral line shapes.

In Reference [25], processes (1) are also considered in the case of an atmosphere of a M red dwarf with an effective temperature $T_{eff} = 3800$ K. The influence of these processes on the population of excited states of the hydrogen atom in this case was investigated using the PHOENIX code [10,16], which as a result generates a model of the considered atmosphere. In the framework of this work, the PHOENIX code includes processes (1) for $n \ge 4$. Some of the results obtained in [25] are illustrated by the Figure 11, where the values of the parameter ζ are presented, i.e., the ratio of the populations of the excited states of the hydrogen atom, calculated with and without taking into account processes (1). The presented figures show that, at least in the region $n \le 15$, processes (1) equally affect the populations of the excited states of the hydrogen atom.

Then, Reference [25] shows that the processes (1) in the whole region of n > 1 also influence the free electron density (see Figure 12). This figure shows the behavior of free electron density calculated with these processes (solid curve) and without them (dashed curve).



Figure 11. (a) the behavior of the population ratio ζ for $3 \le n \le 9$ as a function of the column mass; (b) same as in (a) but for $10 \le n \le 15$ (from [25]).



Figure 12. Structure of model atmosphere–electron density *Ne* and temperature *Te* as a function of column mass.

The presented results suggested that processes (1), due to their influence on the excited state populations and the free electron density, also should influence the atomic spectral line shapes in the atmospheres of late type dwarfs.

2.5. Influences of Chemi-Ionization/Recombination Processes on the Hydrogen Spectral Lines in the M Red Dwarf Atmosphere

In connection with this problem, in Reference [26], the atmospheres of a M red dwarf with an effective temperature $T_{eff} = 3800$ K was also examined. In contrast to the previous case, processes (1) with n = 2 and 3 were included here.

The results presented in [26] show that the chemi-ionization/recombination processes (1), are directly affecting the population of the excited states of the hydrogen atom and the electron concentration, and thus have a very strong effect on the shape of the spectral lines of the atom. For the given atmosphere, the profiles of a number of spectral lines of the hydrogen atom were calculated.

Figure 13 (from [26]) show the line profiles of H_{α} , H_{δ} , H_{ϵ} Pa_{ϵ} with and without inclusion of processes (1). Profiles are synthesized with PHOENIX code with Stark broadening contribution calculated using tables from [40] for Stark broadening of hydrogen lines (linear Stark effect). Lineshape changes, especially in the wings, show the influence of the electron density change having a direct influence on the Stark broadening of hydrogen lines.



Figure 13. Line profiles with (full) and without (red tiny) inclusion of chemi-ionization and chemi-recombination processes for H lines.

3. Symmetric Ion–Atom Processes

It is well known [22] that the chemical composition of the primordial gas consists of electrons and species such as: helium—He, He⁺, He²⁺ and HeH⁺; hydrogen—H, H⁻, H⁺, H⁺₂ and H₂; deuterium—*D*, *D*⁺, H*D*, H*D*⁺ and H*D*⁻; lithium—Li, Li⁺, Li⁻, LiH⁻ and LiH⁺. Evaluation of chemical abundances in the standard Big Bang model are calculated from a set of chemical reactions for the early universe [22]. Among them are very important reactions (2) that involve species like H, H⁺, H⁺₂, He⁺₂, He, He⁺, whose role in the primordial star formation is crucial.

Recently, References [41,42] have pointed out that the photodissociation of the diatomic molecular ion in the symmetric cases (2a) are of astrophysical relevance and could be important in modeling of specific stellar atmosphere layers, and they should be included in some chemical models. The data that involves reactions (2) are also useful in hydrogen and helium theoretical and laboratory plasmas research [43].

3.1. Solar Atmosphere: Visible Wavelength Region

The theoretical investigation of processes (2) started in [44] for the hydrogen symmetric case A = H. Then, in [44,45], the processes (2) were considered in relation to the photosphere and the lower solar chromosphere by characterizing their spectral emissivity $\varepsilon_{ia}(\lambda)$, (the spectral density of the radiation energy, which these processes generate from a unit volume per unit time, into an angle 4π). Their contributions have been collated with concurrent processes (5)–(7) by comparative analysis of their influence using the values of the following parameters: $F_{ei}(\lambda) = \varepsilon_{ia}(\lambda)/\varepsilon_{ei}(\lambda)$, $F_{ea}^{ff}(\lambda) = \varepsilon_{ia}(\lambda)/\varepsilon_{ea}^{fb}(\lambda) = \varepsilon_{ia}(\lambda)/\varepsilon_{ea}^{fb}(\lambda)$, where processes $\varepsilon_{ei}(\lambda)$, $\varepsilon_{ea}^{ff}(\lambda)$, and $\varepsilon_{ea}^{fb}(\lambda)$ are their emissivity.

One of the major results obtained in papers of Mihajlov et al. [44,45] for the visible spectral region of the spectrum are illustrated by the Figure 14 for the considered layer of the solar atmosphere on the basis of standard models [39,46]. It was found that in this region processes (2) give a contribution of 10–12% in comparison with dominant processes (7). This fact alone demonstrated that considered ion–atom radiative processes must be taken into account for solar atmosphere modeling. Later estimates showed, however, that the relative influence of the absorption processes (2) on the solar atmosphere opacity should significantly increase at the transition from the considered wavelength region $\lambda \ge 365$ nm to the region $\lambda_H \le \lambda < 365$ nm, where $\lambda_H = 91.1262$ nm is the wavelength that corresponds to the ionization threshold of the H(1s) atom.



Figure 14. (a) the behaviour of the parameter $F_{ea}^{fb}(\lambda)$ as a functions of height *h* in the solar atmosphere for 350 nm $\leq \lambda \leq 800$ nm; (b) the behaviour of the parameter $F_{ea}^{fb}(\lambda)$ as a function of height *h* of the solar atmosphere for 800 nm $\leq \lambda \leq 1250$ nm (from [44]).

3.2. Solar Atmosphere: UV and VUV Wavelength Region

In the far UV region, the emission channels of processes (2) cease to play a role, while the role of absorption channels in this region grows very rapidly. The total absorption rate coefficient $K_{ia} = K_{ia}^a + K_{ia}^b$ for processes (2a,b) as well as the branch coefficient $X = K_{ia}^b/K_{ia}$ which shows the influence of channels 'a' and 'b' in reaction (2) are presented in Figure 15a,b, respectively. The quantities K_{ia} and X are described in detail in [47] as well as their relations with cross sections. From Figure 15a one can see that K_{ia} strongly depends on temperature and wavelength and, from Figure 15b, it follows that, for lower temperatures, processes (2a) are dominant and importance of processes (2b) increases with the increase of temperature as expected.

In accordance with [47], in the Ultraviolet and Vacuum Ultraviolet regions, the influence of the absorption processes in processes (2) is estimated by the parameter, $F_k(\lambda)$ which is the ratio of the absorption coefficient of these processes and the absorption coefficient determined by the concurrent processes (5)–(7) taken together. The results obtained for the parameters $F_K(h)$ in the wavelength region 92 nm $\leq \lambda \leq 350$ nm are presented by Figure 16.

This figure shows that in the significant part of the considered region of altitudes ($-75 \text{ km} \le h \le 1065 \text{ km}$) the absorption processes (2) together give the contribution that varies from about 10% to about 90% of the contribution of the absorption processes (7), which are considered as the main absorption processes in the solar photosphere. In connection with the other known concurrent absorption processes, it is shown that, in the considered region of altitude, there are significant parts where the symmetric processes completely dominate (see also [48]).

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Figure 15. (a) the total absorption coefficient *Kia* as a function of λ and *T*; (b) the branch coefficient *X* as a function of λ and *T*. Calculations from [47] case *A* = H.



Figure 16. The behaviour of the parameter $F_k(\lambda)$ as a functions of height *h* in the solar atmosphere.

3.3. DB White Dwarf Atmospheres: Visible Wavelength Region

The results from [23], obtained for one Koester model (log g = 8 and $T_{eff} = 12,000$ K), have already provided a more realistic picture of the relative importance of He⁻ (6) and He⁺₂ (2a) total absorption processes, at least in the region $\lambda \leq 300$ nm. In a following paper [24], the relative importance of He⁺₂ and other relevant absorption processes in the region $\lambda \geq 200$ nm was examined for several of Koester's (1980) models ($T_{eff} = 12,000, 14,000, 16,000$ K, log g = 7, 8). It was shown that, in all considered cases, the contribution to opacity of the processes of He⁺₂ molecular ion photodissociation and He + He⁺ collisional absorption charge exchange combined is close to or at least comparable with the contribution of the absorption processes (6) and atomic absorption processes (5).

3.4. DB White Dwarf Atmospheres: UV and VUV Wavelength Region

Let us note in this context that, in the case of white dwarf atmospheres with dominant helium component, among all possible symmetric ion–atom absorbtion processes that are allowed by their composition, only the processes (2a,b) have to be taken into account [23,27].

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Figure 17a presents the total absorption rate coefficient $K_{ia} = K^a_{ia} + K^b_{ia}$ for the processes (2a,b) as well as the branch coefficient $X = K^b_{ia}/K_{ia}$ that shows the influence of channels 'a' and 'b' in reaction (2). The quantities K_{ia} and X are described in detail in [27]. Regarding the importance of the particular channel ('a' and 'b') for the processes (2a,b), conclusions are the same as for the case of hydrogen.



Figure 17. (a) the total absorption coefficient *Kia* as a function of λ and *T*; (b) the branch coefficient *X* as a function of λ and *T*. Data from [27], case *A* = He.

For the calculation of spectral properties, the data from the corresponding DB white-dwarf atmosphere models [49] have been used as well as the data for coefficients given in Figure 17. It was established that the processes (2a,b) significantly influence the opacity of the considered DB white dwarf atmospheres, with an effective temperature $T_{eff} \geq 12,000$ K, which fully justifies their inclusion in one of the models of such atmospheres [50]. However, the same comparison demonstrated also that the dominant role in those atmospheres generally still belongs to the concurrent process) only in some layers of those atmospheres, and only within the part 50 nm $< \lambda < 250$ nm of the far UV and EUV region (see Figure 18). The results obtained in research allow for the possibility of estimating which absorption processes give the main contribution to the opacity in DB white dwarf atmospheres in different spectral regions. Therefore, from [23,27,51,52] results, it follows that the helium absorption processes (2a,b) are dominant in the region 70 nm $\leq \lambda \leq 200$ nm, while, in the region $\lambda \geq 200$ nm, the absorption processes (6) have an important role.

From the presented material, it follows that the considered symmetric ion-atom absorption processes cannot be treated only as one channel among many equal channels with influence on the opacity of the solar atmosphere. Namely, these symmetric processes around the temperature minimum increase the absorption of the EM radiation, so that this absorption becomes almost uniform in the whole solar photosphere


Figure 18. (a) behaviour of the quantity $F_{He} = \kappa_{ia}/(\kappa_{ea} + \kappa_{ei})$ (ratio of the absorption coefficients of processes (2), (5) and (6)) within the atmosphere of a DB white dwarf in the case log g = 8 and $T_{eff} = 12,000$ K; (b) as in (a), but for the case log g = 8 and $T_{eff} = 14,000$ K.

4. Conclusions

All the foregoing shows the undoubted influence of the radiation ion-atom processes and chemi-ionization/recombination processes on the optical properties and on the kinetics of weakly ionized layers of stellar atmospheres, and they should be studied from the spectroscopic aspect. In addition, it can be expected that the reported results will be a sufficient reason for including these processes in the models of stellar atmospheres. The further development of research in a new direction must be connected with the investigations of described processes but in the field of modelling in early Universe chemistry.

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Article



Nonlinear Spectroscopy of Alkali Atoms in Cold Medium of Astrophysical Relevance

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Abstract: The time-dependent population dynamics of hyperfine (HF) sublevels of $n^2 p_{3/2}$ atomic states upon laser excitation in a cold medium of alkali atoms is examined. We demonstrate some peculiarities of the absorption HF multiplet formation in *D*2-line resulting from a long interaction time (~200 µs) interaction between light and Na (n = 3) and Cs (n = 6) atoms in a cold and slow sub-thermal ($T \sim 1$ K) beam. We analytically describe a number of *D*2-line-shape effects that are of interest in spectroscopic studies of cold dusty white dwarfs: broadening by optical pumping, intensity redistribution within components of *D*2-line HF multiplet for partially closed transitions and asymmetry of absorption lines induced by AC Stark shifts for cyclic transitions.

Keywords: spectroscopy; alkali atoms; astrophysics

1. Introduction

Investigation of the fluorescence spectrum of sodium atoms is an important data source for astrophysics and, especially, for understanding the physical processes in cool stars, namely, for brown and white dwarfs. For example, the measured precise atmospheric parameters for shortest period binary white dwarfs confirm the existence of metal-rich envelopes around extremely low-mass white dwarfs and allow us to examine the distribution of the abundance of non-hydrogen elements, including Na, as a function of effective temperature and mass [1]. It is worth emphasizing that we can expect the gravitational wave strain for such systems.

Other interesting astrophysical objects are metal polluted white dwarfs and dusty white dwarfs. It has long been suspected that metal polluted white dwarfs (types DAZ, DBZ and DZ) and white dwarfs with dusty disks can possess planetary systems [2]. Therefore, the spectroscopic observations of sodium atoms in these objects can confirm the validity of this hypothesis. It is in fact known that all dusty white dwarfs show evidence for alkali atoms accretion onto their dusty disk.

In reference [3], the observational constraints on the origin of metals and alkali atoms in cool white dwarfs are discussed. The presence of absorption lines of Na, Mg, Fe etc in the photospheres of

cool hydrogen atmosphere DA-type white dwarfs has been an unexplained problem for a long time. However, in [4], it was shown that the metal abundances in the atmospheres of white dwarfs can be explained by episodic accretion events whenever the white dwarf travels through relatively overdense clouds.

Furthermore, Na 8183.27, 8192.81 Å absorption doublet was discovered in astrophysical systems of white dwarf–main sequence binaries. It allows us to investigate in detail the process of mass transfer interaction in these complex astrophysical systems.

In astrophysics, there is a wide interest in the search for dusty white dwarfs with powerful infrared excess. This excess is produced by orbiting dust disk that contains planetary systems and planetesimals. All these materials get accreted onto the white dwarf and enrich its pure hydrogen or/and helium atmosphere. Studying these heavy elements enriched white dwarfs becomes an effective way to measure directly the bulk compositions of extrasolar planetesimals [5]. Na atoms can be the essential part of these heavy-element clouds; therefore, detailed knowledge of Na emission features can be extremely relevant for investigating such astrophysical problems.

Recently, combined Spitzer and ground-based Korea Microlensing Telescope Network observations identified and precisely measured an Earth-mass planet orbiting ultra cool dwarf [6]. Observations of Na atoms in ultra cool dwarfs can become the effective method for determining planetesimals and Earth-mass planets in dusty disks surrounding the central ultra cool dwarf.

We note as well that, already in 1974, Brown [7] detected the first neutral sodium cloud near Jovian satellite Io (see also [8]). The investigation of these sodium clouds is needed for better understanding of the interaction between Io's atmosphere and Jovian magnetosphere [9] and of processes in the Jovian environment [10].

We are concerned here with important particularities of cold alkali atoms absorption spectra in D2-lines (see Figure 1) that result from a long interaction time of light and matter and, as a consequence, from a strong involvement of optical pumping phenomena within hyperfine (HF) components of the ground and excited states. The described novel theoretical predictions are compared with experimental data obtained in sub-thermal (cold) atomic beams [11,12] operated with two different alkali species, namely Na and Cs. Atomic units are used unless stated otherwise.



Figure 1. Transition strength $\tilde{S}_{F''F'}$ (boxed) and branching Π (circled) factors for the hyperfine components of *D*2-lines for (**a**) cesium (transition $6^2s_{1/2} \rightarrow 6^2p_{3/2}$) and (**b**) sodium (transition $3^2s_{1/2} \rightarrow 3^2p_{3/2}$) atoms. For cyclic (closed) transitions, the factor $\Pi = 1$. The natural lifetime τ and saturation threshold $\Omega_{st} = 1/(\tau\sqrt{2})$ of the resonance states are $\tau = 30.5$ ns and $\Omega_{st} = 3.7$ MHz for Cs while $\tau = 16.4$ ns and $\Omega_{st} = 6.9$ MHz for Na, respectively.

2. Formation of HF Absorption Multiplet for Partially Open Transitions

Absorption spectra of alkali atoms have a multiplet form due to HF structure of atomic states (see Figure 1). There are two different classes of multiplet components corresponding to cyclic

(or closed) and partially open (or non-cyclic) transitions between HF sublevels F' and F'' of the resonant excited (e) and the ground (g) states, respectively. In the partially open case, each spontaneously emitted photon induces two transitions $F' \rightarrow F'' = 2,1$ in Na and $F' \rightarrow F'' = 4,3$ in Cs with probabilities Π (see Figure 1). The dimensionless parameter Π is called the branching ratio. The cyclic transitions correspond to $\Pi = 1$ and they are particularly important in cooling and trapping techniques of neutral atoms, for example, in magneto-optical traps (MOTs) [13].

Excitation of the partially open individual transition $F'' \to F'$ is accompanied by optical pumping phenomenon that is usually associated with redistribution of population within HF sublevels of the ground state because of interaction with resonant light fields [14]. Line-shape effects due to optical pumping (the so-called depletion broadening) in the weak excitation limit and long interaction time τ_{tr} were examined in detail in [15]. A convenient approximation in describing the time-dependent population dynamics of HF sublevels F'', F' is a two-level model. Atoms are excited on the atomic transition by monochromatic laser radiation with frequency ω_L , amplitude A_0 , and detuning $\delta \equiv \omega_L - \omega_{eg}$. If the laser intensity is insufficient to saturate the transition, which corresponds to Rabi frequency Ω ($\Omega = \langle e | A_0 \hat{d} | g \rangle$, where \hat{d} is the atomic dipole operator) being below the natural linewidth Γ_e of the excited state, then the rate of spontaneous transitions per one atom is equal to [14]:

$$\Gamma_g = \frac{\Omega^2 \Gamma_e}{4\delta^2 + \Gamma_e^2}; \quad \Omega \ll \Gamma_e. \tag{1}$$

In a partially open system, each spontaneously emitted photon returns to g-state the Π -portion of the population, while the remaining $(1 - \Pi)$ -portion is transferred to states outside the two-level system. It means the depletion rate Γ_{pum} is equal to $\Gamma_{pum} = (1 - \Pi)\Gamma_g$ and the lower state population n_g of the atom decays exponentially $n_g(\tau) \approx \exp(-\tau/\tau_{pum})$ at the time τ upon transit through the laser beam. The corresponding pumping time is $\tau_{pum} = 1/\Gamma_{pum}$, i.e.,

$$\tau_{pum} = \frac{1}{\Gamma_{pum}} = \frac{1}{1 - \Pi} \frac{4\delta^2 + \Gamma_e^2}{\Omega^2 \Gamma_e}.$$
(2)

In experiments involving atom beams crossed by the laser radiation [11,12], we can identify a transit time τ_{tr} . In such conditions, the total population after the atom beam has crossed the laser radiation, $n_g(\infty)$, becomes $n_g(\infty) \approx \exp(-\tau_{tr}/\tau_{pum})$. Thus, the criterion for the development of optical pumping is $\tau_{tr} > \tau_{pum}$, or in terms of Rabi frequencies

$$\Omega > \Omega_{cr}; \quad \Omega_{cr} = \Gamma_e \sqrt{\frac{1 + 4\delta^2 / \Gamma_e^2}{1 - \Pi} \cdot \frac{\tau_e}{\tau_{tr}}}, \tag{3}$$

where $\tau_e = 1/\Gamma_e$ is the radiative lifetime of the upper e-state. The ratio τ_e/τ_{tr} for cold atomic beams acquires values of $\sim 10^{-4}$ ($\tau_e \approx 20$ ns, $\tau_{tr} \approx 200 \ \mu$ s) [3]. Therefore, the depletion manifestation can be observed at Rabi frequencies well below (by two orders of magnitude) the saturation frequency $\Omega_{st} \equiv \Gamma_e/\sqrt{2}$; exceeding this value results in the development of nonlinear (power broadening) and quantum optics effects [14].

Important spectroscopic features of optical pumping emerge when the Rabi frequency is set close, or above, the critical value Ω_{cr} (3) [15]. Conventional experiments would usually register the fluorescence signal *J* (absorption line) from the entire excitation volume, which is proportional to the total number $\Gamma_e \bar{n}_e$ of photons emitted by a single atom. On the other hand, the number $(1 - \Pi)\Gamma_e \bar{n}_e$ of spontaneously emitted photons on transitions outside the *g-e* system is equal to the total loss $1 - n_g(\infty)$ of the ground state population, so that the signal *J* can be written as

$$J \equiv \Gamma_e \bar{n}_e = \frac{1}{(1 - \Pi)} \left(1 - \exp(-\tau_{tr}/\tau_{pum}) \right).$$
(4)

Equation (4) yields a dependence of the fluorescence signal on the laser detuning δ in the following explicit form:

$$J(\delta) = \frac{\Omega^2 \tau_{tr}}{\Gamma_e} \frac{1}{P_{pum}} \left[1 - \exp\left(-\frac{P_{pum}}{1 + 4\delta^2 / \Gamma_e^2}\right) \right],\tag{5}$$

$$P_{pum} = \frac{\tau_{tr}}{\tau_{pum}^0}; \quad \tau_{pum}^0 = \frac{1}{1 - \Pi} \frac{\Gamma_e}{\Omega^2}.$$
 (6)

The above equation and data presented in Figure 2 show that the absorption spectrum strongly depends on the pumping parameter P_{pum} defined as in Equation (6), which is given by the ratio of transit time τ_{tr} and pumping time τ_{pum}^0 . The latter has the meaning of optical pumping time at resonant excitation, when $\delta = 0$. Importantly, the parameter P_{pum} can be large even at laser intensities well below the saturation limit: $P_{pum} >> 1$ when $\tau_{tr} >> \tau_{pum}^0$ and $\Omega << \Omega_{cr}$.



Figure 2. Dependence of the absorption signal *J* Equation (5) on the reduced detuning δ/Γ_e for different values of the pumping parameter P_{pum} Equation (6) (shown as labels of the curves). All curves are unity-normalized at the line center $\delta = 0$.

According to Equation (4), optical pumping leads to saturation of the fluorescence signal at the value of $1/(1 - \Pi)$. This fact along with redistribution of intensities within D2-line multiplet have been predicted and experimentally demonstrated in [15]. Another phenomenon described by Equations (4) and (5) is related to depletion broadening of the absorption line. As an estimation of the characteristic full width Δ_{pm} of the spectral absorption profile induced by optical pumping, one can write

$$\Delta_{pm} = \sqrt{\frac{\tau_{tr}}{\tau_e} (1 - \Pi) \cdot \Omega^2 - \Gamma_e^2}.$$
(7)

In the case of cold beams of alkali atoms, with $\tau_{tr}/\tau_e \sim 10^4$ and $\Omega > \Omega_{cr}$ for $\delta = 0$ (see Equation (3)), the width Δ_{pm} acquires the form $\Delta_{pm} \approx \Omega \sqrt{(1-\Pi)\tau_{tr}/\tau_e} \sim 100 \cdot \Omega$, which may essentially exceed the power broadening effects ($\Delta_{pw} \sim \Omega$) even in the limit of weak excitation. The corresponding experimental investigation of spectral broadening due to optical pumping (the parameter $\tau_{tr}/\tau_e \sim 100$) in partially open HF level systems in Na has been reported in [15]. We note that, since the experiments were performed at low number densities ($\sim 10^{10} \text{ cm}^{-3}$) of Na atoms, line-shape modifications by radiation trapping in collimated beams [16] can be disregarded.

3. Experimental Setup and Spectroscopic Data

Common experimental practice foresees the use of two types of thermal beams: effusive beam with longitudinal velocity $v_{lg} \approx 400 \text{ m/s}$ [17], and crossed beams with $v_{lg} \approx 600 \text{ m/s}$ [18]. The second type represents a supersonic beam with $v_{lg} \approx 1100 \text{ m/s}$ [15] and a rather specific velocity distribution function [19]. An additional experimental possibility is represented by laser cooled atomic beams [20,21], which are characterized by small v_{lg} around 12 m/s, which corresponds to the sub-thermal temperature interval lying below 1 K and practically negligible transversal velocity.

At the core of the relevant setup is a hollow pyramid with reflective inner surfaces and a hole at its vertex. By shining a single, large diameter (35 mm) laser beam along the pyramid axis, the optical configuration of an MOT is achieved. Imbalance of the radiation pressure along the pyramid axis pushes Cs atoms out of the pyramid hole. Further collimation of the atoms is achieved by a transverse optical molasses right after the pyramid hole (Figure 3). The atom beam is then excited by a diode laser tunable over the hyperfine transitions belonging to the *D*2-line of Cs; blue laser radiation is superposed in order to ionize the excited atoms. The production rate of ions is proportional to the total number of excited atoms in the excited population [12]. Note that line-shape modifications by radiation trapping in a cold medium [22,23] can be disregarded for the above experimental conditions.



Figure 3. Schematic diagram of the experimental setup for the production and excitation of a cold Cs atom beam.

Figure 4 shows an experimentally obtained absorption profile (dots) upon excitation of the cyclic $F'' = 4 \rightarrow F' = 5$ transition for Cs atoms. We underline two main points: (i) although the main line-profile results from symmetric power broadening; (ii) there is a slight asymmetry in the line-shape.

This asymmetry is induced by the other HF components of $n^2 p_{3/2}$ sublevels and, as it will be shown in the next section, is strongly affected by the relationships between laser Rabi frequency and values of HF splitting. For comparison, we show in Figure 5 a situation occurring for Na atoms in similar experimental conditions to demonstrate the occurrence of a quite non-standard line shape, whose explanation is presented in the following.



Figure 4. Theoretical (solid curve, Equation (19)) and experimental [11] (dots) absorption profile of the *D*2-line of Cs upon an excitation of the closed $F'' = 4 \rightarrow F' = 5$ transition in a cold sub-thermal Cs beam for laser power of 1.0 mW (the corresponding Rabi frequency $\Omega = 21.2$ MHz). The dashed curve corresponds to the power broadening profile. The bar-dashed curve exhibits the natural broadening profile.



Figure 5. The same as in Figure 4 in the case of *D*2-line of sub-thermal Na atoms and the excitation of the closed $F'' = 2 \rightarrow F' = 3$ transition with laser Rabi frequency $\Omega = 21.2$ MHz.

4. Cyclic Transitions Treatment: Modeling and Discussion

The description of cyclic transitions can rarely be reduced to the two-level system model: even upon resonant excitation ($\delta = 0$) in a cold beam, the presence of other HF levels may result in the appearance of fundamentally new effects. Let us add the third level $|3\rangle$ to a two-level closed system $|4\rangle$, $|2\rangle$ (see Figure 6a), which can decay to the passive state $|1\rangle$ due to spontaneous emission.

We associate states $|1\rangle$, $|2\rangle$ and $|3\rangle$, $|4\rangle$ of Figure 6 with HF g-sublevels F'' = 1, F'' = 2 and e-sublevels F' = 2, F' = 3, accordingly, in the case of Na atoms (see Figure 1), while, for Cs HF structure, we choose HF g-sublevels F'' = 3, F'' = 4 and e-sublevels F' = 4. F' = 5.



Figure 6. (a) schematic illustration of a three-level system having a cyclic transition $|4\rangle \rightarrow |2\rangle$ and a partially open transition $|3\rangle \rightarrow |2\rangle$ with the branching factor Π . The additional passive state $|1\rangle$ accumulates the population due to the spontaneous transition $|3\rangle \rightarrow |1\rangle$; (b) the same scheme in the rotating wave approximation. The bare ground state $|2\rangle$ is selected as the position of zero energy (dashed horizontal line).

We are concerned here with the resonant excitation of cyclic HF transitions between HF sublevels. As a consequence, we set the following constrains for laser detuning $\delta = \omega_L - \omega_{42}$, as indicated in Figure 6: $\delta = -\Delta_4 \sim \Gamma \ll \Delta_3 \approx \Delta_{34}$, where $\Gamma = \Gamma_3 = \Gamma_4$ is the unique natural linewidth of the upper $n^2 p_{3/2}$ levels. The lower levels $|1\rangle$, $|2\rangle$ correspond to the HF sublevels of the ground state $n^2 s_{3/2}$. Its large HF splitting (Δ_1), compared to the one (Δ_{34}) of resonant states $n^2 p_{3/2}$, transforms the state $|1\rangle$ into a dark state that is not interacting with the pump laser. Figure 6b shows the dressed state configuration obtained using the rotating wave approximation [14]. If the energy ε_2 of the state 2 is chosen as zero, then the energies $\varepsilon_{3,4}$ of the dressed states $|3\rangle$, $|4\rangle$ turn out to be determined by the laser detuning $\delta: \varepsilon_4 = \Delta_4 = -\delta$, $\varepsilon_3 = -\Delta_3$.

It is convenient to represent the Rabi frequencies $\Omega_{F''F'}$ of the pump laser in terms of a single reduced frequency Ω_{red} [15], which is the product of the amplitude A_0 of the laser field and the reduced dipole matrix element $D_{S,P} = (n^2 S_{1/2} ||D|| n^2 P_{3/2})$ [24] for the respective non HF resolved fine transition: $\Omega_{red} = A_0 D_{S,P}$. The partial Rabi frequency values $\Omega_{F''F'}$ for HF transitions $S_{1/2}, F'' \rightarrow P_{3/2}, F'$ are determined by the corresponding, so-called, line strengths $S_{F''F'}$ [24]:

$$\Omega_{F''F'} = \Omega_{red} \sqrt{\tilde{S}_{F''F'}}; \quad \tilde{S}_{F''F'} = S_{F''F'} / D_{S,P}^2.$$
(8)

The values of non-dimensional parameters $\tilde{S}_{F''F'}$ are reported within rectangular frames in Figure 1. In Figure 6b, and subsequent discussions, we are using the abbreviations $\Omega_i = \Omega_{2''i'}$ (*i* = 3, 4).

A fundamentally new aspect for the three-level system is the appearance of the dynamic (AC) Stark shifts [13,14] of the state 2, due to its laser induced mixing with the state 3. As a result, the energy defect (detuning) between states 4 and 2 is also undergoing a shift (see also below, Equation (16))

$$\Delta_4 \to \delta_4 = \Delta_4 - \Omega_3^2 / (4\Delta_{34}). \tag{9}$$

In the case of a resonance ($\Delta_4 = 0$), the increase of the pump laser intensity leads to an increasing shift of the actual detuning δ_4 . It is worth noting that, in order to significantly affect the light-induced

asymmetry in the line profile, the absolute value $|\delta_4|$ of detuning must be larger than the natural linewidth Γ , i.e., according to Equation (9), it is necessary that Ω_3 exceeds the saturation threshold: $\Omega_3^2 > 2\Gamma\Delta_3 > \Gamma^2$.

In the weak excitation limit ($\Omega_4 < \Gamma$), the induced transitions $|2\rangle \rightarrow |4\rangle$ and $|2\rangle \rightarrow |3\rangle$ are independent from each other, as they are relatively weak compared to spontaneous transitions. This means that, due to the partially open $|2\rangle \rightarrow |3\rangle$ transition, the ground state depletion should take place and, consequently, the cyclic transition $|2\rangle \rightarrow |4\rangle$ should be affected by the depletion broadening. The results of Section 2 are applicable for the partially open transition $|2\rangle \rightarrow |3\rangle$. In particular, using the notation of Figures 1 and 3, the relation (2) can be rewritten as

$$\frac{\tau_{pum}}{\tau_{tr}} \approx \frac{1}{\tau_{tr}\Gamma_{pum}} = \frac{\tau/\tau_{tr}}{1-\Pi} \frac{4\Delta_3^2}{\Omega_2^2}.$$
(10)

The inequality $\tau_{tr}/\tau_{pum} > 1$ characterizing the onset of optical pumping is fulfilled at $\Omega_3 > \Omega_{cr} = 1.50$ MHz and $\Omega_3 > \Omega_{cr} = 9.6$ MHz for Na and Cs, respectively, for the experimental ratio $\tau_{tr}/\tau \sim 10^4$. It is well seen that, in the case of Cs, its critical value Ω_{cr} lies beyond the saturation threshold. At such Rabi frequencies, the linear approximation is no longer applicable and a more accurate approach is required to describe the light–matter interaction.

4.1. Adiabatic Approach

The large value of the ratio $\tau_{tr}/\tau \sim 10^4$ allows one to use the method of adiabatic elimination [14,25] to obtain explicit qualitative description of the above effects for cyclic transitions. The exact analysis of the dynamics of optical pumping should be carried out within the framework of the density matrix [14] for the three-level system model shown in Figure 6b. Equations describing the interaction of a single atom (from the atomic beam) with a classical exciting radiation are the optical Bloch Equations [14]:

$$i\dot{\rho}_{ij} = \omega_{ij}\rho_{ij} + \sum_{k} \left(H_{ik} \ \rho_{kj} - \rho_{ik} \ H_{kj}\right) + il_{ij},\tag{11}$$

where non-diagonal elements ρ_{ij} ($i \neq j$) are associated with the so-called coherence while diagonal elements give the level population $n_i:n_i = \rho_{ii}$. The matrix l_{ij} describes relaxation processes due to spontaneous radiative transitions. The matrix H_{ik} determines the interaction of atoms with the laser light. If the electric field of the excitation laser in Figure 3 has a Gaussian distribution, $E(z) = A_0 \exp(-z^2/(2d^2))$, with width *d* along the atomic beam axis (coordinate *z*), then in the rotating wave approximation (RWA) [14] H_{ik} has the following representation (see also Equation (8)):

$$\Omega_{F''F'}(t) = 0.5 \cdot \Omega_{red} \sqrt{\tilde{S}_{F''F'}} \exp(-2t^2/\tau_{tr}^2).$$
(12)

The time of flight of an atom through the excitation zone is determined by the laser beam waist size *d*: $\tau_{tr} = 2 d/v$. Under the experimental conditions of [12], the respective values are $\tau_{tr} \approx 1.1$ mm and $\tau_{tr} \approx 200 \ \mu$ s.

It is worth noting that all individual members of Equation (12) may be rewritten in the universal form:

$$\dot{X}_k = -\left(\Gamma_k + i\omega_k\right)X_k + F_k(t). \tag{13}$$

Equation (13) can be interpreted as the equation of a linear oscillator with complex coordinates X_k , which are subject to a slowly varying external force $F_k(t)$. One can see that the frequency detuning ω_k plays the role of rigidity, while the width Γ_k is associated with the dissipation constant. From classical mechanics, it is well known that characteristic time τ_{rel} of the evolution of a forced oscillator

to a steady-state condition is determined as $\tau_{rel} = 1/\sqrt{\Gamma_k^2 + \omega_k^2}$ [26]. If $\tau_{tr} >> \tau_{rel}$, the left-hand side of Equation (13) becomes negligible, so that Equation (13) has the following solutions:

$$X_k = F_k(t) / (\Gamma_k + i\omega_k); \tau_{tr} >> 1 / \sqrt{\Gamma_k^2 + \omega_k^2}.$$
(14)

4.2. Strict Results

Adiabatic elimination for Bloch equations (11) was implemented in [27]. At the first stage, using Equation (14), we express non-diagonal elements ρ_{ij} ($i \neq j$) via diagonal ones, i.e., via populations n_i . As a result, the following closed equation systems describe the population dynamics:

$$\dot{n}_2 = \Gamma n_4 + \Gamma \Pi n_3 + r_4 (n_4 - n_2) + r_3 (n_3 - n_2), \tag{15a}$$

$$\dot{n}_i = -\Gamma n_i - r_i (n_i - n_2); \quad i = 3, 4,$$
 (15b)

$$r_{i} = \Omega_{i2}^{2} \frac{\Gamma_{i}}{4\delta_{i}^{2} + \Gamma_{i}^{2}}, \quad \delta_{i} = \omega_{i2} - \frac{\Omega_{j2}^{2}}{4\omega_{43}}, \quad \Gamma_{i} = \Gamma\left(1 + \frac{\Omega_{j2}^{2}}{2\omega_{43}^{2}}\right), \quad j \neq i, 2,$$
(16)

where ω_{43} is the difference between energies ε_4 and ε_3 . These relations have the structure of balance equations for population transfer between the levels due to the spontaneous, with rate Γ , and laser stimulated, with rate r_i , transitions. An important result is the appearance of the AC Stark shifts $\Omega_{i2}^2/(4\omega_{34})$ (see also Equation (9)) in the pumping rate constant r_i .

Owing to the adiabatic elimination, system (15) can be further reduced to a single equation for the total population N(t) ($N = n_2 + n_4 + n_3$) [25] of a single atom:

$$\dot{N} = -\frac{\Gamma \cdot (1-\Pi) \cdot r_3}{\Gamma + 2r_3 + r_4 \left(\Gamma + r_3\right) / \left(\Gamma + r_4\right)} N,$$
(17)

$$n_2 = \frac{N}{1 + r_4 / (\Gamma + r_4) + r_3 / (\Gamma + r_3)}; \quad n_i = \frac{r_i}{\Gamma + r_i} n_2 \quad i = 3, 4.$$
(18)

This reduction allows us to obtain an explicit representation for the fluorescence signal *J*, which is proportional to the total number $\Gamma(\bar{n}_4 + \bar{n}_3)$ of photons emitted by a single atom from the excitation volume

$$J = J(t)|_{t=\infty} \equiv \Gamma \int_{-\infty}^{t} d\tau \cdot (n_4(\tau) + n_3(\tau))|_{t=\infty}.$$
 (19)

4.3. Discussion of the Line-Shape Structure

Solid lines in Figures 4 and 5 are plotted using Formula (19). Note that the theoretical results well describe the experimental data. Both profiles for Na and Cs atoms have an asymmetry, manifested in a rapid drop of the right wings. This asymmetry is due to the upwards AC Stark energy shift (see Equations (9) and (16)) of level $| 2 \rangle$ in Figure 6, which corresponds to the HF sublevel F'' = 2 for Na and F'' = 4 for Cs. As a result, the value of actual detuning $\delta_4 = \Delta_4 - \Omega_3^2 / \Delta_{43}$ depends on the sign of $\Delta_4 = -\delta$, i.e., the actual detuning δ_4 for the same absolute value of laser detuning δ is larger for the right wing side compared to the left wing side.

There is another point that complicates the situation. The structure of Equation (17) implies a slow decay of the population resulting from a weak laser stimulated mixture between states $|2\rangle$ and $|3\rangle$. It means the cyclic transition $|2\rangle \rightarrow |3\rangle$ ceases to be closed. The decay factor should manifest itself in a depletion broadening. The actual profile, thus, is formed via interplay of space dependent (see Equations (16) and (17)) optical pumping and AC Stark effects.

5. Conclusions

One of the characteristic features of a cold medium is the long interaction time τ_{tr} between light and atoms. As a result, a variety of nonlinear optical effects may take place even for moderate values of light power. We have experimentally observed and theoretically modeled such nonlinear effects by studying HF-selective laser interaction with cold atomic beams consisting of alkali atoms.

Significant modifications of the optical features are found in closed transitions. In particular, we have predicted and experimentally demonstrated the appearance of an asymmetry in the corresponding absorption lines and have explained this occurrence through AC Stark shifts of the involved states. The long transit time (~0.2 ms) through the excitation zone, combined with a relatively small mixing of HF sublevels of the resonant $n^2 p_{3/2}$ state due to the laser coupling results in opening a decay channel for cyclic transitions. The particularities of the line shape formation are a result of the strong interplay between time dependent optical pumping and AC Stark effects.

The results discussed here are of potential interest for interpretation of spectroscopic data obtained from fluorescence spectra of a cold medium of astrophysical relevance such as different modifications of cold white dwarfs or neutral sodium clouds near Jovian moon Io.

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Article





Radiative and Collisional Molecular Data and Virtual Laboratory Astrophysics

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Abstract: Spectroscopy has been crucial for our understanding of physical and chemical phenomena. The interpretation of interstellar line spectra with radiative transfer calculations usually requires two kinds of molecular input data: spectroscopic data (such as energy levels, statistical weights, transition probabilities, etc.) and collision data. This contribution describes how such data are collected, stored, and which limitations exist. Also, here we summarize challenges of atomic/molecular databases and point out our experiences, problems, etc., which we are faced with. We present overview of future developments and needs in the areas of radiative transfer and molecular data.

Keywords: atomic/molecular data; radiative and collisional processes; stars

1. Introduction

Many fields in astronomy such as astrophysics, astrochemistry and astrobiology, depend on data for atomic and molecular (A + M) collision and radiative processes. Among these amount of data collections there are atomic and molecular processes and spectral regions that even today are poorly represented. Therefore, there is an urgent need to collect these data in the databases as well as to develop methods for improving the existing ones. Also, this require a joint effort both of scientists and IT software specialists to develop state-of-the-art infrastructures satisfying their needs, such as Virtual Laboratory [1–3].

The Base Astrophysical Targets

Nowadays, the data in the field of astrophysics modeling are especially important and needed for simulations/calculations. For example the A + M data for hydrogen are important for development of atmosphere models of solar and near solar type stars and for radiative transport investigations as well as an understanding of the kinetics of stellar and other astrophysical plasmas [4,5]. Modern codes for stellar atmosphere modelling, like e.g., PHOENIX (see e.g., [6–8]) require the knowledge of atomic data, so that the access to such atomic data, via online databases become very important.

The helium A + M data are of interest particularly for helium-rich white dwarf atmospheres investigations [9,10]. Such data are also important in modelling early Universe chemistry (see the paper of Coppola et al., 2013 [11]). The data for H and some metal atoms like Li, Na, Si are important for the exploring of the geo-cosmical plasmas, the interstellar medium as well as for studies of the early Universe chemistry and for the modelling of stellar and solar atmospheres (see, e.g., [12,13]).

Recently, in the papers [14–18] it has been pointed out that the photodissociation of the diatomic molecular ion in the symmetric and non-symmetric cases, are of astrophysical relevance and could be important in modeling of specific stellar atmosphere layers and they should be included in some chemical models. In the symmetric case, it was considered the processes of molecular ion photodissociation (bound-free) and ion-atom photoassociation (free-bound):

$$h\nu + A_2^+ \Longleftrightarrow A + A^+, \tag{1}$$

where *A* and A^+ are atom and ion in their ground states, and A_2^+ is molecular-ion in the ground electronic state.

In the non-symmetric case, the similar processes of photodissociation/photoassociation are:

$$h\nu + AM^+ \iff A^+ + M,$$
 (2)

where M is an atom whose ionization potential is less than the corresponding value for atom A. AM^+ is also molecular-ion in the ground electronic state.

In the general case molecular ion A_2^+ or AM^+ can be in one of the states from the group which contains the ground electronic state. For the solar atmosphere, *A* usually denotes atom H(1s) and *M* one of the relevant metal atoms (Mg, Si, Ca, Na) [14–16], but there are cases where *A* = He, and *M* = H, Mg, Si, Ca, Na. For the helium-rich white dwarf atmospheres *A* denotes He(1s²) and *M* denotes, H(1s), and eventually carbon or oxygen [19,20].

Recently, the results from [16] show the importance of including the symmetric processes with A = H(1s) in the stellar atmosphere models like [5]. Also, for modeling the DB white dwarf atmospheres results for case $A = He(1s^2)$ have been used (Koester 2016, private communication). The photodissociation of HeH⁺ has been extensively studied both from a theoretical and experimentalist point of view and inserted in chemical networks describing the formation and destruction of primordial molecules.

It is well known [21] that the chemical composition of the primordial gas consists of electrons and species such as: helium- He, He⁺, He²⁺ and HeH⁺; hydrogen- H, H⁻, H⁺, H₂⁺ and H₂; deuterium- *D*, D^+ , HD, HD⁺ and HD⁻; lithium- Li, Li⁺, Li⁻, LiH⁻ and LiH⁺. Evaluation of chemical abundances in the standard BB model are calculated from a set of chemical reactions for the early universe [21] and is presented at Figure 1 from [21]. One can see that among them are species like molecular ions H₂⁺, HD⁺, HeH⁺, etc., whose role in the primordial star formation is important.



Figure 1. Evaluation of chemical abundances in the standard Big Bang (BB) model. Vertical axes are the relative abundances and the horizontal axes are relative to the redshift (taken from [21]).

2. MolD

2.1. Database Description

MolD database contains cross-sections for photodissociation processes [22], as well as corresponding data on molecular species and molecular state characterisations. MoID project is part of Serbian Virtual Observatory (SerVO)¹ and Virtual Atomic and Molecular Data Center (VAMDC)² (see Figure 2).

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Figure 2. Snapshot from the query page from the Virtual Atomic and Molecular Data Center (VAMDC) portal.

MolD application is implemented as a customisation and extension of NodeSoftware provided by VAMDC. It complies to VAMDC interoperability standards and protocols for distributed remote queries. The underlying technology is Python-based, with Django as a Web framework [23], MySQL as a relational database system [24]. The web application runs on Apache web server.

The data model of MolD application is tailored to specifically suit the needs of the theoretical photodissociation data, and yet to easily map onto VAMDC's standardized XSAMS³ (XML Schema for Atoms, Molecules and Solids format) schema for representation and exchange of atomic and molecular data⁴.

2.2. Accessing MolD Data

MolD data can be accessed in several ways:

Via MolD homepage (http://servo.aob.rs/mold). There is an AJAX-enabled (Asynchronous JavaScript and XML) web form for data querying as well as calculating and plotting average thermal cross sections along available wavelengths for a given temperatiure.

¹ http://servo.aob.rs

² http://vamdc.eu

http://vamdc-standards.readthedocs.io/en/latest/dataModel/vamdcxsams/structure.html
 http://standards.vamdc.eu

- Via VAMDC portal (http://portal.vamdc.eu), where one can pose a distributed query to 30 databases across the European scientific institutes.
- Via standalone applications which support VAMDC-TAP (Table Access Protoco) for data access and tranformation to VAMDC-XSAMS.

During 2017, MolD entered *stage* 3 of development. Currently, the database includes cross-section data for processes which involve species such as He_2^+ , H_2^+ , MgH^+ , HeH^+ , LiH^+ , NaH^+ . These processes are important for exploring of the interstellar medium, the early Universe chemistry as well as the modeling of different stellar and solar atmospheres (see papers [11,15,16,20,22]).

Our plans include transition to new versions of Django framework and NodeSoftware, with ongoing incremental inclusion of A + M data from our papers. We also intend to develop a more intuitive interface for querying and presentation of multidimensional data on our website.

2.3. Example: H_2^+ Molecular Ion

MolD is available online from the end of 2014 and it contains the data for the photodissociation processes Equation (1) with A = H(1s) and $A = He(1s^2)$. Also it contains the relevant data for some other non-symmetric photodissociation processes Equation (2) where M = Li, Na, Mg or He.

The methods of calculation. The cross-sections for the photodissociation of individual ro-vibrational state of the considered molecular ion H_2^+ is determined in the dipole approximation [14]:

$$\sigma_{J,v}(\lambda) = \frac{8\pi^3}{3\lambda} \left[\frac{(J+1)|D_{E,J+1;v,J}|^2 + J|D_{E,J-1;v,J}|^2}{2J+1} \right],\tag{3}$$

and the corresponding averaged thermal cross sections are given by:

$$\sigma_{\rm phd}(\lambda,T) = \frac{1}{Z} \sum_{J} \sum_{v} g_{J;v} (2J+1) e^{-\frac{E_{Jv} - E_{00}}{k_{\rm B}T}} \sigma_{J,v}(\lambda).$$
(4)

where *T* is temperature, λ -wavelength, $D_{E,J+1;v,J}$ is the radial matrix element [25], E_{Jv} is the energy of the individual states with the angular and vibrational quantum numbers *J* and *v* respectively, and *Z* is the partition function

$$Z = \sum_{J} \sum_{v} g_{J,v} (2J+1) e^{-\frac{E_{Jv} - E_{0,0}}{k_{\rm B}T}}.$$
(5)

In this expression the product $g_{J,v} \times (2J + 1)$ is the statistical weight of the considered state and the coefficient $g_{J,v}$ depends on the "the spin of the nuclei".

The photo-dissociation crosssection $\sigma_{phd}(\lambda, T)$ given by Equation (4), as well as the coefficients $K_{ia}(\lambda, T)$, are determined within the approximation where the processes are treated as the result of the radiative transitions between the ground and the first excited adiabatic electronic state of the molecular ion H₂⁺ which are caused by the interaction of the electron component of the ion-atom system with the free electromagnetic field taken in the dipole approximation.

For determination of $\sigma_{phd}(\lambda, T)$, as well as the coefficients $K_{ia}(\lambda, T)$ it is important to know the dipole matrix element $D_{12}(R)$ defined by relations

$$D_{12}(R) = \mathbf{D}_{12}(\mathbf{R}), \quad \mathbf{D}_{12}(\mathbf{R}) = \langle 1 | \mathbf{D}(\mathbf{R}) | 2 \rangle \tag{6}$$

where $R = |\mathbf{R}|$ and $\mathbf{D}(\mathbf{R})$ is the operator of electron dipole momentum. The mentioned adiabatic electronic states, $X^2 \sum_g^+$ and $A^2 \sum_u^-$, are denoted here with $|1\rangle$ and $|2\rangle$ and R is the internuclear distance in the considered ion-atom system.

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The described mechanism of the processes causes absorption of the photon with energy ϵ_{λ} near the resonant point $R = R_{\lambda}$, where R_{λ} is the root of the equation

$$U_{12}(R) \equiv U_1(R) - U_2(R) = \epsilon_{\lambda}.$$
(7)

where $U_1(R)$ corresponds to the ground electronic state, and $U_2(R)$ —to the first excited electronic state.

In Figure 3 are presented the data for $\sigma_{J,v}(\lambda)$ Equation (3) for the case J = 0 and v = 10, in the wavelength region 50 nm $\leq \lambda \leq 1500$ nm.



Figure 3. The behaviour of the cross-section $\sigma_{J,v}(\lambda)$ Equation (3) for J = 0 and v = 10, as a function of λ .

The spectral coefficients. The absorption process (1) i.e., processes of molecular ion photodissociation (bound-free) for the case A = H is characterized by partial spectral absorption coefficients $\kappa_{ia}(\lambda)$ (see e.g., [25]) taken in the form

$$\kappa_{ia}(\lambda) = \sigma_{phd}(\lambda, T) N(\mathrm{H}_2^+) \tag{8}$$

where $N(\text{H}_2^+)$ is density of the H_2^+ and $\sigma_{phd}(\lambda, T)$ is average cross-section for photo-dissociation of this molecular ion given by Equation (4). As in previous papers [14,25] the partial spectral absorption coefficient $\kappa_{ia}(\lambda)$ is also used in the form

$$\kappa_{ia}(\lambda) = K_{ia}(\lambda, T)N(H)N(H^{+})$$
⁽⁹⁾

where the coefficient $K_{ia}(\lambda, T)$ is connected with $\sigma_{phd}(\lambda, T)$ by the relations

$$K_{ia}(\lambda, T) = \sigma_{phd}(\lambda, T)\chi^{-1}, \quad \chi = \frac{N(H)N(H^+)}{N(H_2^+)}.$$
 (10)

In accordance with the definition of the absorption coefficient $\kappa_{ia}(\lambda)$, the coefficient $K_{ia}(\lambda, T)$ is given in units (cm⁵). The results for the average photodissociation cross-section $\sigma_{phd}(\lambda, T)$ for H_2^+ molecular ion are illustrated by Figure 4. The curves in this figure show the behavior of $\sigma_{phd}(\lambda, T)$ as a function of λ for a wide range of temperatures *T*, which are relevant for the stelar atmosphere (e.g., solar photosphere). The values of the coefficient $K_{ia}(\lambda, T)$, defined by Equation (10), are presented in the Table 1 for the regions 90 nm $\leq \lambda \leq$ 370 nm with small wavelength steps and for 3000 K $\leq T \leq$ 10,000 K in order to enable easier use (interpolation) of this results. This allows direct calculation of the spectral absorption coefficients during the process of applying a any atmosphere model with the given parameters of plasma and composition.



Figure 4. The behaviour of the averaged cross-section $\sigma_{phd}(\lambda, T)$ for photodissociation of the H₂⁺ molecular ion, as a function of λ and *T*.

λ (nm)	3000 K	4000 K	5000 K	6000 K	7000 K	8000 K	9000 K	10,000 K
90	3.50E-37	2.62E-38	5.50E-39	1.93E-39	9.04E-40	5.07E-40	3.20E-40	2.20E-40
91	3.75E-37	2.78E-38	5.80E-39	2.03E-39	9.47E - 40	5.30E - 40	3.34E - 40	2.29E - 40
92	3.99E-37	2.94E-38	6.10E-39	2.12E-39	9.90E-40	5.53E - 40	3.48E - 40	2.39E - 40
93	4.23E-37	3.10E-38	6.40E-39	2.22E-39	1.03E-39	5.76E - 40	3.62E - 40	2.48E - 40
94	4.47E - 37	3.25E-38	6.69E-39	2.31E-39	1.07E-39	5.98E - 40	3.76E - 40	2.57E - 40
95	4.71E - 37	3.41E-38	6.98E-39	2.41E-39	1.12E-39	6.20E-40	3.89E-40	2.66E - 40
96	4.95E - 37	3.56E-38	7.27E-39	2.50E-39	1.16E-39	6.42E - 40	4.03E - 40	2.75E - 40
97	5.18E-37	3.70E-38	7.54E-39	2.59E-39	1.20E-39	6.63E-40	4.16E - 40	2.84E - 40
98	5.40E - 37	3.85E-38	7.82E-39	2.68E-39	1.24E-39	6.85E - 40	4.29E - 40	2.92E - 40
99	5.62E-37	3.99E-38	8.08E-39	2.76E-39	1.27E-39	7.05E - 40	4.41E - 40	3.01E - 40
100	5.82E-37	4.12E - 38	8.34E-39	2.85E-39	1.31E-39	7.26E - 40	4.54E - 40	3.09E - 40
101	6.03E-37	4.25E - 38	8.59E-39	2.93E-39	1.35E-39	7.46E - 40	4.66E - 40	3.17E - 40
102	6.22E-37	4.38E-38	8.83E-39	3.01E-39	1.38E-39	7.65E - 40	4.78E - 40	3.26E - 40
103	6.40E - 37	4.50E - 38	9.06E-39	3.09E-39	1.42E - 39	7.84E - 40	4.90E - 40	3.34E - 40
104	6.58E-37	4.62E - 38	9.29E-39	3.16E-39	1.45E - 39	8.03E - 40	5.02E - 40	3.41E - 40
105	6.74E - 37	4.73E - 38	9.51E-39	3.24E-39	1.49E - 39	8.21E - 40	5.13E - 40	3.49E - 40
106	6.90E-37	4.83E - 38	9.72E-39	3.31E-39	1.52E - 39	8.39E - 40	5.24E - 40	3.57E - 40
107	7.04E - 37	4.93E - 38	9.92E-39	3.38E-39	1.55E - 39	8.57E - 40	5.35E - 40	3.64E - 40
108	7.18E-37	5.03E - 38	1.01E - 38	3.44E - 39	1.58E - 39	8.74E - 40	5.46E - 40	3.72E - 40
109	7.30E-37	5.11E - 38	1.03E - 38	3.51E-39	1.61E - 39	8.90E - 40	5.56E - 40	3.79E - 40
110	7.41E - 37	5.20E-38	1.05E - 38	3.57E-39	1.64E - 39	9.06E - 40	5.67E - 40	3.86E - 40
111	7.52E-37	5.28E - 38	1.06E - 38	3.62E-39	1.67E - 39	9.22E - 40	5.77E - 40	3.93E - 40
112	7.61E-37	5.35E - 38	1.08E - 38	3.68E-39	1.69E-39	9.37E - 40	5.86E - 40	3.99E - 40
113	7.69E-37	5.42E - 38	1.09E - 38	3.73E-39	1.72E-39	9.52E - 40	5.96E - 40	4.06E - 40
114	7.77E-37	5.48E - 38	1.11E - 38	3.79E-39	1.74E - 39	9.66E - 40	6.05E - 40	4.12E - 40
115	7.83E-37	5.54E - 38	1.12E - 38	3.83E-39	1.77E - 39	9.80E - 40	6.14E - 40	4.18E - 40
116	7.89E-37	5.59E-38	1.13E-38	3.88E-39	1.79E-39	9.93E-40	6.22E - 40	4.24E - 40

Table 1. The coefficient K_{ia} (cm⁵) Equation (9) as a function of λ and T.

Table 1. Cont.

λ (nm)	3000 K	4000 K	5000 K	6000 K	7000 K	8000 K	9000 K	10,000 K
117	7.93E-37	5.63E-38	1.15E-38	3.93E-39	1.81E-39	1.01E-39	6.31E-40	4.30E-40
118	7.97E-37	5.68E-38	1.16E - 38	3.97E-39	1.83E-39	1.02E - 39	6.39E-40	4.36E - 40
119	8.00E-37	5.71E - 38	1.17E - 38	4.01E-39	1.85E-39	1.03E-39	6.46E - 40	4.42E - 40
120	8.02E-37	5.75E-38	1.18E-38	4.05E - 39	1.87E-39	1.04E - 39	6.54E - 40	4.47E - 40
121	8.03E-37	5.78E-38	1.18E - 38	4.08E-39	1.89E-39	1.05E - 39	6.62E - 40	4.52E - 40
122	8.04E-37	5.80E-38	1.19E - 38	4.12E-39	1.91E-39	1.06E-39	6.69E - 40	4.57E - 40
123	8.04E - 37	5.82E - 38	1.20E - 38	4.15E - 39	1.93E-39	1.07E - 39	6.76E - 40	4.62E - 40
124	8.03E-37	5.84E - 38	1.21E - 38	4.18E - 39	1.94E - 39	1.08E - 39	6.83E - 40	4.67E - 40
125	8.01E - 37	5.85E - 38	1.21E - 38	4.21E - 39	1.96E - 39	1.09E - 39	6.89E - 40	4.72E - 40
126	7.99E-37	5.86E - 38	1.22E - 38	4.23E-39	1.97E - 39	1.10E - 39	6.96E - 40	4.77E - 40
127	7.97E-37	5.87E - 38	1.22E - 38	4.26E - 39	1.99E-39	1.11E - 39	7.02E - 40	4.81E - 40
128	7.94E - 37	5.87E - 38	1.23E - 38	4.28E - 39	2.00E-39	1.12E - 39	7.08E - 40	4.86E - 40
129	7.90E-37	5.87E - 38	1.23E - 38	4.30E - 39	2.01E - 39	1.13E - 39	7.14E - 40	4.90E - 40
130	7.86E-37	5.87E - 38	1.23E - 38	4.32E - 39	2.03E-39	1.14E - 39	7.19E - 40	4.94E - 40
131	7.81E - 37	5.86E - 38	1.24E - 38	4.34E - 39	2.04E - 39	1.15E - 39	7.25E - 40	4.98E - 40
132	7.76E-37	5.86E - 38	1.24E - 38	4.36E - 39	2.05E - 39	1.15E - 39	7.30E - 40	5.02E - 40
133	7.70E-37	5.84E - 38	1.24E - 38	4.37E - 39	2.06E - 39	1.16E - 39	7.35E - 40	5.06E - 40
134	7.65E - 37	5.83E - 38	1.24E - 38	4.39E-39	2.07E - 39	1.17E - 39	7.40E - 40	5.10E - 40
135	7.58E - 37	5.82E - 38	1.24E - 38	4.40E - 39	2.08E - 39	1.17E - 39	7.45E - 40	5.13E - 40
136	7.52E - 37	5.80E - 38	1.24E - 38	4.41E - 39	2.09E-39	1.18E - 39	7.49E - 40	5.17E - 40
137	7.45E - 37	5.78E - 38	1.24E - 38	4.42E - 39	2.09E-39	1.19E - 39	7.54E - 40	5.20E - 40
138	7.38E-37	5.76E - 38	1.24E - 38	4.43E - 39	2.10E - 39	1.19E - 39	7.58E - 40	5.24E - 40
139	7.31E-37	5.73E - 38	1.24E - 38	4.43E - 39	2.11E - 39	1.20E - 39	7.62E - 40	5.27E - 40
140	7.24E - 37	5.71E - 38	1.24E - 38	4.44E - 39	2.12E - 39	1.20E - 39	7.66E - 40	5.30E - 40
141	7.16E - 37	5.68E - 38	1.24E - 38	4.45E - 39	2.12E - 39	1.21E - 39	7.70E - 40	5.33E - 40
142	7.08E - 37	5.65E - 38	1.24E - 38	4.45E - 39	2.13E-39	1.21E - 39	7.74E - 40	5.36E - 40
143	7.00E - 37	5.62E - 38	1.23E - 38	4.45E - 39	2.13E-39	1.22E - 39	7.77E - 40	5.39E - 40
144	6.92E - 37	5.59E - 38	1.23E - 38	4.46E - 39	2.14E - 39	1.22E - 39	7.80E - 40	5.41E - 40
145	6.84E - 37	5.56E - 38	1.23E - 38	4.46E - 39	2.14E - 39	1.22E - 39	7.84E - 40	5.44E - 40
146	6.76E-37	5.52E - 38	1.23E - 38	4.46E - 39	2.15E - 39	1.23E - 39	7.87E - 40	5.46E - 40
147	6.68E-37	5.49E - 38	1.22E - 38	4.46E - 39	2.15E - 39	1.23E - 39	7.90E - 40	5.49E - 40
148	6.59E-37	5.45E - 38	1.22E - 38	4.46E-39	2.15E-39	1.23E-39	7.93E - 40	5.51E - 40
149	6.51E-37	5.42E-38	1.22E-38	4.45E-39	2.15E-39	1.24E-39	7.95E-40	5.54E - 40
150	6.42E-37	5.38E-38	1.21E-38	4.45E-39	2.16E-39	1.24E-39	7.98E-40	5.56E - 40
151	6.34E-37	5.34E-38	1.21E-38	4.45E-39	2.16E-39	1.24E-39	8.01E - 40	5.58E-40
152	6.25E-37	5.30E-38	1.20E-38	4.44E-39	2.16E-39	1.25E-39	8.03E-40	5.60E-40
153	6.17E-37	5.26E-38	1.20E-38	4.44E-39	2.16E-39	1.25E-39	8.05E-40	5.62E - 40
154	6.08E-37	5.22E-38	1.20E-38	4.43E-39	2.16E - 39	1.25E-39	8.08E-40	5.64E - 40
155	6.00E-37	5.16E-36	1.19E-38	4.43E-39	2.16E-39	1.25E-39	8.10E-40	5.66E - 40
150	5.91E-57	5.14E-38	1.19E-30	4.42E-39	2.17E-39	1.25E - 39	8.12E - 40	5.68E-40
157	5.63E-37	5.10E-38	1.16E-36	4.42E-39	2.17E-39	1.26E-39	8.14E-40	5.70E-40
150	5.74E-57	5.00E-30	1.10E-30	4.41E-39	2.17E-39	1.20E-39	0.10E-40	5.72E-40
159	5.00E-37	5.02E-38	1.17E-38	4.40E-39	2.17E-39	1.26E-39	8.18E-40	5.73E-40
160	5.36E-37	4.90E-30	1.17E-30	4.39E-39	2.17E-39	1.20E-39	8.19E-40	5.73E-40
162	5.41E-37	4.90E-38	1.10L-38	4.37E-39	2.10E-39	1.20E-39	8.22E-40	5.78E-40
164	5.35E-37	4.05E-50	1.13E-38	4.30E-39	2.10E-39	1.20E-39	8.24E-40	5.79E - 40 5.81E 40
165	5.25E - 37 5.17E - 37	4.01E - 38 4.77E - 38	1.14E = 38 1.14E = 38	4.33E-39	2.10E - 39 2.16E - 39	1.27E - 39 1.27E - 39	8.23E - 40 8.27E - 40	5.81E - 40 5.82E - 40
166	5.09E - 37	4.77E-38	1.14E - 30 1.13E - 38	4.34E-39	2.10E - 39 2.16E - 39	1.27E - 39 1 27E - 39	8.28E - 40	5.02E - 40 5.83E - 40
167	5.02E - 37 5.02E - 37	4.68E-38	1.13E - 38	4.32E-39	2.10E - 39	1.27E - 39	8 29E - 40	5.85E - 40
168	4.94E - 37	4.60E - 38	1.13E - 38 1.12E - 38	4.32E - 39 4.31E - 39	2.10E - 39 2.16E - 39	1.27E - 39 1.27E - 39	8.30E - 40	5.86E - 40
169	4.86E - 37	4.60E - 38	1.12E - 38	4.30E - 39	2.15E - 39	1.27E - 39	8.31E-40	5.87E - 40
170	4.79E - 37	4.56E - 38	1.11E - 38	4.29E - 39	2.15E - 39	1.27E - 39	8.32E - 40	5.88E - 40
171	4.71E - 37	4.52E - 38	1.10E - 38	4.27E - 39	2.15E - 39	1.27E - 39	8.33E-40	5.89E - 40
172	4.64E - 37	4.48E - 38	1.10E - 38	4.26E - 39	2.15E - 39	1.27E - 39	8.34E - 40	5.90E - 40
173	4.57E - 37	4.43E-38	1.09E - 38	4.25E - 39	2.14E-39	1.27E - 39	8.34E - 40	5.91E - 40
174	4.50E - 37	4.39E-38	1.09E - 38	4.24E - 39	2.14E-39	1.27E - 39	8.35E-40	5.92E - 40
175	4.43E-37	4.35E-38	1.08E-38	4.22E-39	2.14E-39	1.27E-39	8.36E-40	5.93E-40

Table 1. Cont.

λ (nm)	3000 K	4000 K	5000 K	6000 K	7000 K	8000 K	9000 K	10,000 K
176	4.36E-37	4.31E-38	1.07E-38	4.21E-39	2.13E-39	1.27E-39	8.36E-40	5.94E-40
177	4.29E-37	4.27E - 38	1.07E - 38	4.20E-39	2.13E-39	1.27E-39	8.37E - 40	5.94E - 40
178	4.22E-37	4.23E-38	1.06E - 38	4.18E-39	2.13E-39	1.27E-39	8.37E - 40	5.95E - 40
179	4.15E - 37	4.19E - 38	1.06E - 38	4.17E-39	2.12E-39	1.27E-39	8.38E - 40	5.96E - 40
180	4.09E-37	4.15E - 38	1.05E - 38	4.16E-39	2.12E-39	1.27E-39	8.38E - 40	5.96E - 40
181	4.02E-37	4.11E - 38	1.04E - 38	4.14E-39	2.12E-39	1.27E-39	8.38E - 40	5.97E - 40
182	3.96E-37	4.07E - 38	1.04E - 38	4.13E-39	2.11E-39	1.26E-39	8.39E-40	5.98E - 40
183	3.90E-37	4.03E-38	1.03E-38	4.11E-39	2.11E-39	1.26E-39	8.39E-40	5.98E - 40
184	3.84E-37	4.00E - 38	1.03E-38	4.10E-39	2.11E-39	1.26E-39	8.39E-40	5.99E - 40
185	3.78E-37	3.96E-38	1.02E - 38	4.09E-39	2.10E-39	1.26E-39	8.39E-40	5.99E - 40
186	3.72E-37	3.92E-38	1.01E - 38	4.07E-39	2.10E-39	1.26E-39	8.39E-40	6.00E - 40
187	3.66E-37	3.88E-38	1.01E - 38	4.06E-39	2.09E-39	1.26E-39	8.39E-40	6.00E-40
188	3.60E-37	3.84E-38	1.00E - 38	4.04E - 39	2.09E-39	1.26E-39	8.40E - 40	6.01E - 40
189	3.54E-37	3.81E-38	9.95E-39	4.03E-39	2.09E-39	1.26E-39	8.40E - 40	6.01E - 40
190	3.49E-37	3.77E-38	9.89E-39	4.01E-39	2.08E-39	1.26E-39	8.40E - 40	6.02E-40
191	3.43E-37	3.73E-38	9.83E-39	4.00E-39	2.08E-39	1.26E-39	8.39E - 40	6.02E - 40
192	3.38E-37	3.70E-38	9.77E-39	3.98E-39	2.07E-39	1.26E-39	8.39E-40	6.02E-40
193	3.33E-37	3.66E-38	9.71E-39	3.97E-39	2.07E-39	1.25E-39	8.39E - 40	6.03E-40
194	3.27E-37	3.63E-38	9.65E-39	3.95E-39	2.06E-39	1.25E-39	8.39E - 40	6.03E-40
195	3.22E-37	3.59E-38	9.59E-39	3.94E-39	2.06E-39	1.25E-39	8.39E-40	6.03E-40
196	3.17E-37	3.56E-38	9.54E-39	3.92E-39	2.06E-39	1.25E-39	8.39E-40	6.03E-40
197	3.12E-37	3.52E-38	9.48E-39	3.91E-39	2.05E-39	1.25E-39	8.38E - 40	6.04E - 40
198	3.08E-37	3.49E-38	9.42E-39	3.89E-39	2.05E-39	1.25E-39	8.38E - 40	6.04E - 40
199	3.03E-37	3.46E-38	9.36E-39	3.88E-39	2.04E-39	1.25E-39	8.38E - 40	6.04E - 40
200	2.98E-37	3.42E-38	9.31E-39	3.86E-39	2.04E-39	1.24E - 39	8.38E - 40	6.04E - 40
205	2.76E-37	3.26E-38	9.03E-39	3.79E-39	2.01E-39	1.24E - 39	8.35E - 40	6.04E - 40
210	2.56E-37	3.11E-38	8.75E-39	3.71E-39	1.99E-39	1.23E-39	8.33E-40	6.04E - 40
215	2.37E-37	2.97E-38	8.49E-39	3.64E-39	1.96E-39	1.22E-39	8.29E-40	6.03E-40
220	2.20E-37	2.83E-38	8.24E-39	3.57E-39	1.94E-39	1.21E-39	8.25E - 40	6.02E - 40
225	2.04E - 37	2.71E-38	7.99E-39	3.50E-39	1.91E-39	1.20E-39	8.21E - 40	6.01E - 40
230	1.90E-37	2.59E-38	7.76E-39	3.43E-39	1.89E-39	1.19E-39	8.17E - 40	5.99E - 40
235	1.77E - 37	2.48E-38	7.53E-39	3.36E-39	1.86E-39	1.18E-39	8.13E - 40	5.98E - 40
240	1.66E - 37	2.37E-38	7.32E-39	3.30E-39	1.84E - 39	1.17E - 39	8.08E - 40	5.96E - 40
245	1.55E - 37	2.27E-38	7.11E-39	3.23E-39	1.81E - 39	1.16E - 39	8.04E - 40	5.94E - 40
250	1.45E - 37	2.18E - 38	6.92E-39	3.17E-39	1.79E-39	1.15E - 39	7.99E - 40	5.92E - 40
255	1.36E - 37	2.09E-38	6.73E-39	3.11E-39	1.76E - 39	1.13E-39	7.94E - 40	5.90E - 40
260	1.27E - 37	2.01E - 38	6.54E - 39	3.05E-39	1.74E - 39	1.12E - 39	7.89E - 40	5.87E - 40
265	1.20E - 37	1.93E - 38	6.37E-39	2.99E-39	1.72E - 39	1.11E - 39	7.83E - 40	5.84E - 40
270	1.12E - 37	1.85E - 38	6.20E-39	2.94E - 39	1.69E-39	1.10E - 39	7.78E - 40	5.82E - 40
275	1.06E - 37	1.78E - 38	6.04E - 39	2.89E-39	1.67E - 39	1.09E - 39	7.72E - 40	5.79E - 40
280	9.99E-38	1.72E - 38	5.89E-39	2.83E-39	1.65E - 39	1.08E - 39	7.67E - 40	5.76E - 40
285	9.43E-38	1.66E - 38	5.74E - 39	2.78E-39	1.63E-39	1.07E - 39	7.62E - 40	5.73E - 40
290	8.92E-38	1.60E - 38	5.60E-39	2.73E-39	1.61E - 39	1.06E - 39	7.56E - 40	5.70E - 40
295	8.45E - 38	1.54E - 38	5.47E - 39	2.69E-39	1.59E - 39	1.05E - 39	7.51E - 40	5.67E - 40
300	8.00E-38	1.49E - 38	5.34E - 39	2.64E - 39	1.57E - 39	1.04E - 39	7.46E - 40	5.64E - 40
305	7.60E-38	1.44E - 38	5.22E-39	2.60E-39	1.55E - 39	1.03E - 39	7.41E - 40	5.61E - 40
310	7.22E-38	1.39E - 38	5.10E - 39	2.56E-39	1.53E - 39	1.02E - 39	7.36E - 40	5.59E - 40
315	6.86E-38	1.35E - 38	4.99E - 39	2.52E-39	1.51E - 39	1.01E - 39	7.32E - 40	5.56E - 40
320	6.53E-38	1.31E - 38	4.88E - 39	2.48E-39	1.49E - 39	1.01E - 39	7.27E - 40	5.53E - 40
325	6.23E-38	1.27E - 38	4.78E-39	2.44E-39	1.48E - 39	9.96E-40	7.22E - 40	5.51E - 40
330	5.94E - 38	1.23E - 38	4.68E-39	2.40E-39	1.46E - 39	9.88E-40	7.17E - 40	5.48E - 40
335	5.67E - 38	1.19E - 38	4.58E - 39	2.37E-39	1.44E - 39	9.79E - 40	7.13E - 40	5.45E - 40
340	5.42E - 38	1.16E - 38	4.49E - 39	2.33E-39	1.43E - 39	9.71E - 40	7.08E - 40	5.43E - 40
345	5.19E - 38	1.13E - 38	4.40E - 39	2.30E-39	1.41E - 39	9.63E - 40	7.04E - 40	5.40E - 40

Table 1. Cont.

λ (nm)	3000 K	4000 K	5000 K	6000 K	7000 K	8000 K	9000 K	10,000 K
350	4.97E-38	1.09E-38	4.31E-39	2.26E-39	1.40E-39	9.55E-40	6.99E-40	5.37E-40
355	4.76E - 38	1.06E - 38	4.23E-39	2.23E-39	1.38E-39	9.47E - 40	6.95E - 40	5.35E - 40
360	4.57E - 38	1.04E - 38	4.15E - 39	2.20E-39	1.37E - 39	9.39E - 40	6.90E - 40	5.32E - 40
365	4.38E - 38	1.01E - 38	4.07E - 39	2.17E-39	1.35E - 39	9.32E - 40	6.86E - 40	5.29E - 40
370	4.21E - 38	9.82E-39	4.00E-39	2.14E-39	1.34E - 39	9.24E - 40	6.81E - 40	5.26E - 40

Solar atmosphere: absorption processes. The influence of radiation processes (1) can be estimated by comparing their intensities with the intensities of known concurrent radiation processes, namely:

$$\varepsilon_{\lambda} + \begin{cases} H^{-} \\ H + e^{'} \end{cases} \Rightarrow H + e^{''}$$
(11)

$$\varepsilon_{\lambda} + \begin{cases} H^* \\ H^+ + e' \end{cases} \Rightarrow H^+ + e''$$
(12)

The relative contributions of the processes (1), with respect to processes (11) and (12), is described by the quantities F_{κ} defined by relation

$$F_{\kappa} = \frac{\kappa_{ia}}{\kappa_{ea} + \kappa_{ei}} \tag{13}$$

where κ_{ea} is absorption spectral coefficient of processes (11) and κ_{ei} is absorption spectral coefficient of processes (12) (see papers [14,16]).

Similarly to the He case in DB white dwarf atmosphere [25] calculations of the absorption coefficient were performed for the solar photosphere and lower chromosphere by means of a standard Solar atmosphere model [5], and the total contribution of the processes (1) to the solar opacity was estimated [16]. The results of the calculations of the parameter F_{κ} for 92 nm $\leq \lambda \leq$ 350 nm are presented in Figure 5. The figure show that in the significant part of the considered region of altitudes ($-75 \text{ km} \leq h \leq 1065 \text{ km}$) the absorption process (1) together give the contribution which varies from about 10% to about 90% of the contribution of the absorption process (11) and (12), which are considered as the main absorption processes [16].



Figure 5. Upper panel: The behavior of the temperature *T*, *N*_{*H*} and *Ne* as a function of height *h* within the considered part of the solar atmosphere model; lower panel: A surface plot of the quantity $F_{\kappa} = \kappa_{ia}/(\kappa_{ea} + \kappa_{ei})$ (data taken from [14]) as a function of λ and height *h* for a model of the solar photosphere [5].

On the basis of the above, it can be concluded that photodissociation processes represent important channels for destruction of molecules in lot of astrophysical environments and features of the interacting radiation are important in their spectral analyses.

3. Future Developments and Concluding Remarks

Exploitation the full potential of A + M data and database services is an ongoing challenge in virtual data centers. There are still many limitations and problems that users are facing with such as poor documentation, missing of data evaluation, no open access, etc. The aim of MolD database is to be accessible, and be used by the wider scientific community, through VAMDC and to follow certain protocols and defined rules in order to eliminate such limitations and problems.

The next step of development i.e., the stage three of MolD development will be the implementation of possibility to fit the tabulated data. We plan to develop fitting formulas for photodissociation cross section as the function of the temperature and wavelength. Also, we intend to update the current database with newly calculated/measured data.

The continuation of such developments and services such as constantly updated online A + M database, is crucial in the field of astrophysics and modern physics due to its rapid development and make an immense impact on the way science is done in the developing world.

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Article



Rosetta Mission: Electron Scattering Cross Sections—Data Needs and Coverage in BEAMDB Database

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Abstract: The emission of [O I] lines in the coma of Comet 67P/Churyumov-Gerasimenko during the Rosetta mission have been explained by electron impact dissociation of water rather than the process of photodissociation. This is the direct evidence for the role of electron induced processing has been seen on such a body. Analysis of other emission features is handicapped by a lack of detailed knowledge of electron impact cross sections which highlights the need for a broad range of electron scattering data from the molecular systems detected on the comet. In this paper, we present an overview of the needs for electron scattering data relevant for the understanding of observations in coma, the tenuous atmosphere and on the surface of 67P/Churyumov-Gerasimenko during the Rosetta mission. The relevant observations for elucidating the role of electrons come from optical spectra, particle analysis using the ion and electron sensors and mass spectrometry measurements. To model these processes electron impact data should be collated and reviewed in an electron scattering database and an example is given in the BEAMD, which is a part of a larger consortium of Virtual Atomic and Molecular Data Centre—VAMDC.

Keywords: electron scattering; cross sections; Rosetta mission; atomic and molecular databases

1. Introduction

The Rosetta spacecraft was launched in 2004 as a part of the European Space Agency (ESA) space program, with the mission to rendez-vous with, orbit and place a lander upon periodic comet 67P/Churyumov-Gerasimenko. Rosetta was in orbit with the cometary nucleus from 2014 to September 2016 during which time it was able to closely examine how the coma of the comet and the frozen comet's surface changed relative to distance from the Sun. On November 2014 Rosetta dispatched a lander, Philae, which touched down on the comet's surface and recorded, for the first time, in situ data from the surface. This pioneering mission has provided us with new and unexpected data that are changing our understanding of the structure and chemistry of cometary systems and their role in the evolution of our solar system and possible origins of life on Earth. For example, the D to H ratio in cometary water ice is very different from that on Earth and, among the other similar findings, challenges the hypothesis that water on Earth was brought by cometary impact [1].

In this paper, we will review another intriguing and unexpected result from the Rosetta mission namely the role of electron induced dissociation in the comet's coma. The data needed to model electron

processes in cometary coma and its possible relevance to the formation but also the dissociation and fragmentation of molecules observed by Rosetta instruments, will be discussed together with the current data available and the databases in which such data may be found.

2. Rosetta Instruments and Their Observation of Electron Scattering Processes in the Cometary Coma

Rosetta orbiter carried eleven different complex scientific instruments, while the Philae lander had ten instruments. Only those that are immediately relevant for the case study of the role of electrons in comas and the detection of more complex species that may be formed by electron induced chemistry will be reviewed here.

2.1. FUV Emissions Measured by the ALICE Instrument

ALICE was a far-ultraviolet (FUV) imaging spectrograph that could specially resolve spectra in the range from 70 to 205 nm. Coma emission and the reflected solar spectrum from the nucleus were recorded using ALICE throughout the Rosetta encounter. The coma was identified by a spectrum that contains several features that are weak in the solar spectrum and do not appear in the reflected light from the nucleus [2]. Besides strong hydrogen Lyman lines, lines from oxygen multiplets at 98.9, 115.2, 130.4 and 135.6 nm were observed, as were weak multiplets from carbon C I lines at 156.1 and 165.7 nm and emission bands coming from CO. The surprise was the O I line at 135.6 nm, originated from the forbidden transition ${}^{5}S^{o}_{-3}P$ since this is usually not seen in comas. The presence of this line and the intensity ratio of H I and O I multiplets is characteristic of the process of electron dissociative excitation of water molecules [3] and led Feldman et al. [2] to establish that electron collisions with H₂O is the dominant source of these emissions. Similarly, they attributed C I emissions to electron dissociative excitation of CO₂. The relative contribution of the UV and electron impact to the dissociation processes are dependent on the location with respect to the nucleus and the heliocentric distance as discussed in [4,5].

2.2. Observations from the OSIRIS Instrument

OSIRIS (Optical, Spectroscopic and Infrared Remote Imaging System) was one of Rosetta's major imaging systems equipped with a wide-angle camera (WAC) and a series of narrow band filters covering range from 245 to 640 nm and two broad band filters, green and red, covering the spectra up to 720 nm. This instrument recorded coma emission lines and specifically targeted to the transitions of O, OH, CN, CS, NH and NH₂. The mapping of water distribution was possible indirectly through observations of O I and OH bandpass filters. The O I filter covers the forbidden transitions from the O I ($2p^4$) ¹D state which is populated directly by photodissociation of H₂O molecules, while the OH filter covers the (0–0) band of the A $^{2}\Sigma^{+}$ –X $^{2}\Pi$ transition of OH, centred at about 308.5 nm, which is excited almost entirely by fluorescence of sunlight as pointed out in [6]. The O I ¹D state can be also populated from the transition from the O I ($2p^4$) ^{1}S state. Within the CN filter lies an emission line B $^{2}\Sigma^{+}$ –X $^{2}S^{+}$ (0, 0) at 388 nm, within the NH₂ filter there is a wide emission band $\tilde{A} \, ^{2}A_{1}$ –X $^{2}B_{1}$ (0, 10, 0) and the NH filter covers the NH A $^{3}\Pi_{1}$ –X $^{3}\Sigma^{-}$ (0–0) transition [6].

From this data Bodewits et al. [6] derived column densities and calculated global production rates using the standard Haser model. They found that the water production rates derived from OH are larger than those derived from [O I], OH and [O I] photolysis. Indeed, they analysed all production rates and found a much larger drop in water production rates than diurnal variation can explain. Therefore, they concluded that the photo-dissociation and fluorescence could solely explain the processes resulting in the OH, [O I], CN and NH emission observed in the inner coma and that the fragments might emanate from different parent species and/or be formed by other processes [6]. One additional process is electron induced dissociation of water, when including this in their model a much better fit was obtained, indeed electron driven dissociation of water was found to be dominant in agreement with Feldman et al. [2].

2.3. Detection of Organic Molecules on the Comet Surface—COSAC Mass Spectrometry

The COSAC (COmetary SAmpling and Composition) experiment and Ptolemy were two gas analysers on the Philae lander built to monitor the chemical composition of the surface of comet 67P. Due to difficulties in landing Philae was not able to deploy all of its instruments as planned and the drill could not be deployed to collect samples for in-situ analysis. However, seven measurements were made by both COSAC and Ptolemy during Philae's hopping and at its final landing site in a so-called 'sniff mode' that had no active sampling but rather just ionized whatever molecules were present in the ionization chamber of the mass spectrometer. The sample with the richest data was acquired a few minutes after the first touchdown with subsequent decay of signal strength in the other six measurements. Both instruments measured a nearly identical decay of both the water (m/z 18) and CO (m/z 28) peaks. However, in the COSAC measurements the peak at m/z 44 decays much slower than all the other ion species, including the water peak and, the m/z 44 peak also decays much slower in the COSAC measurements than in the Ptolemy data. From these results, it has been concluded that COSAC analysed a regolith sample from the cometary nucleus in situ while Ptolemy measured cometary gas from the ambient coma [7].

The compounds detected by COSAC are listed in Table 1. All of the larger molecules can be formed from the smaller compounds carbon monoxide (CO), methane (CH₄), water (H₂O) and ammonia (NH₃) by simple addition reactions [8]. The m/z 44 peak measured by COSAC was likely dominated by organic species, e.g., from acetaldehyde (C₂H₄O), formamide (HCONH₂) and acetamide (CH₃CONH₂), whereas the peak measured by Ptolemy was interpreted to be mostly due to CO₂. Recently, a comparison and comparative analysis of the Rosetta mass spectrometers (COSAC/Ptolemy/ROSINA) that puts some question mark on the presence of some of the nitrogen-bearing species was presented [9]. Ptolemy measurements confirmed many of the species observed by COSAC and through observation of regular peaks in the observed mass distributions indicated the presence of a sequence of compounds with additional -CH₂- and -O- groups (mass/charge ratios 14 and 16, respectively) which confirms COSAC's observations of acetaldehyde and may be explained by the presence of a radiation-induced polymer at the surface. Ptolemy measurements also indicated an apparent absence of aromatic compounds such as benzene and neither H₂S nor SO₂ were observed [10]. Ammonia believed to be the precursor of N containing compounds was not unambiguously detected by either Ptolemy or COSAC, probably due to its tendency to adsorb on stainless steel surface.

Name of Compound	Sum Formula	Abundance wrt Water
Methane	CH ₄	0.5%
Water	H ₂ O	100%
Hydrogen cyanide	HCN	0.9%
Carbon monoxide	CO	1.2%
Methylamine	CH ₃ NH ₂	0.6%
Acetonitrile	CH ₃ CN	0.3%
Isocyanic acid	HNCO	0.3%
Acetaldehyde	CH ₃ CHO	0.5%
Formamide	HCONH ₂	1.8%
Ethylamine	$C_2H_5NH_2$	0.3%
Methyl isocyanate	CH ₃ NCO	1.3%
Acetone	CH ₃ COCH ₃	0.3%
Propionaldehyde	C ₂ H ₅ CHO	0.1%
Acetamide	CH ₃ CONH ₂	0.7%
Glycolaldehyde	CH ₂ OHCHO	0.4%
Ethylene glycol	HOC ₂ H ₄ OH	0.2%

 Table 1. List of molecules identified on the comet nucleus of comet 67P by the COSAC instrument [8].

 Abundances are given normalized to water, which is the most abundant compound.

Recent experiments on the irradiation of ice mixtures reveal that many of the larger molecules can be formed by electron bombardment, often at low energies and this will be discussed further in Section 5. Indeed, the bombardment and dissociation of ice species has been proposed as a route by which molecular oxygen can form. One of the most unexpected observations of Rosetta through the Rosina instrument was the detection of molecular oxygen as the fourth most abundant gas in the atmosphere of comet 67P. Oxygen is reactive so it was felt that it is unlikely to survive long periods in space. The amount of molecular oxygen detected showed a strong relationship to the amount of water measured at any given time, suggesting that their origin on the nucleus and release mechanism are linked and that irradiation of water ice leading to oxygen production and storage in the ice is a plausible mechanism for oxygen formation on a comet [11].

2.4. Electrons in the Cometary Coma

That there are copious amounts of electrons to induce such dissociative excitation was confirmed by ion and electron sensors (RPC-IES) on the Rosetta craft. Concentrations of particles and their time evolution in inner coma plasma was measured by Rosetta Plasma Consortium (RPC) [12] using a set of sensors developed for this purpose. The Ion and Electron Sensor (IES) was an electrostatic plasma analyser that covered an energy/charge range from 1 eV/e to 22 keV/e with a resolution of 4% [13]. The sensor provided 3D ion and electron distributions over the whole measured energy range. It was capable of simultaneously measuring electrons and positive ions with the single entrance aperture owing to two back-to-back top-hat geometry analysers. The LAP instrument (Langmuir probes) measured the plasma density in the range of $(10^0-10^6 \text{ cm}^{-3})$, electron temperature (10^2-10^5 K) and plasma flow velocity (up to 10^4 ms^{-1}). The LAP also measured the AC electric field up to 8 kHz [14]. The LAP was complemented by the Mutual Impedance Probe, MIP which probes the plasma and measured the natural plasma frequency which yields the electron density in the range from 2 cm⁻³ to $1.5 \times 10^5 \text{ cm}^{-3}$ and temperature from 30 K to 10^6 K [15].

Depending on the comet distance from the Sun, both the solar wind and solar radiation interact with its nucleus and inner coma shielded by comet's own atmosphere and ionosphere. Cometary ions are created by photoionization of neutral species, mainly like H₂O and CO₂ and their products from photodissociation, and by charge transfer with solar wind protons [16]. Solar wind electrons in interplanetary space typically have Maxwellian distribution functions with thermal energies of several eV to tens of eV [13]. This energy distribution of electrons differs from one created by photoionization of cometary neutrals by solar radiation in the cometary comas at certain distances from the Sun. Electrons of cometary origin are mainly the product of photochemistry, originating from direct photoionization and from Auger processes. They are thermalized by collisions, elastic and inelastic.

The electron density in the coma was measured complementary by the RPC Langmuir Probe (LAP) and Mutual Impedance Probe (MIP). The first findings of the spatial distribution of the plasma near comet 67P/CG showed a highly structured pattern that indicated an origin from local ionization of neutral gas. The electron density fell off with distance as 1/r in the range from 8 km from the nucleus up to 260 km [17]. Edberg et al. [17] concluded that this is in accord with a model in which the ionization of a neutral gas is expanding radially from the comet nucleus and when there is no significant recombination or other loss source for the plasma. However, they warned that the observed data have a large scatter around fitted 1/r curve and that results could be an average effect of combination of transport electric fields and solar wind.

From such data, the suprathermal energy distribution of electrons could be derived. The electron energy distribution near the comet depends on the comet distance from perihelion and mass loading process when the atoms and molecules in the cometary coma are photoionized and then interact with the solar wind flow. It spans from the energy distribution of the solar wind itself to the modified distribution where electrons are significantly decelerated as a consequence of magnetic field causing regions to pile-up. The mass loading process is connected to the outgassing rate of the comet. The formation of suprathermal electrons which are accelerated from a few eV upward

to hundreds of eV, thus can play an important role in the electron driven chemistry of the comet. The Rosetta IES sensor recorded the presence of suprathermal electrons at larger distances than expected from the previous models of such weakly outgassing comets [18]. The observed electron energy distributions change by reducing a heliocentric distance from pure solar wind distribution to non-Maxwellian one that include suprathermal electrons showing maxima at energies from 10 eV to 300 eV [18]. Clark et al. [18] hypothesize that the most likely mechanisms of creating accelerated electron distributions are heating by waves generated by the pick-up ion instability and by the mixing of cometary photoelectrons, secondaries and solar wind electrons.

Further statistical analysis of Rosetta IES sensor recordings by performing fitting procedures that involve two separate sub-populations of electrons below and above 8.6 eV mean energy [19] revealed different relationships between their density and temperature and possible mechanisms of creating suprathermal electrons. Broiles et al. [19] suggested that electrons above 8.6 eV are being heated by waves driven by counter streaming solar wind protons. This conclusion arises from the observations that the population of electrons above 8.6 eV correlates well with the density of local neutrals, while the sub-population below 8.6 eV is dominated by the local magnetic field strength. Recently, Deca et al. [20] have used a fully 3D kinetic model to simulate the ion and electron dynamics of the solar wind interaction with a weakly outgassing comet 67P. They used a detailed kinetic treatment of the electron dynamics in order to cover energy distribution of electrons and to identify the origin of the warm and suprathermal electrons.

Electron energy ranges that correspond to the relevant processes in electron collisions with atoms and molecules are shown in Figure 1. The elastic cross section is dominating over low electron energies and usually is prominent even at higher energies where the ionization cross section becomes comparable in magnitude. Vibrational excitations are important at low energies but they also extend to higher energies due to resonance decay. Attachment and dissociation processes are relevant in certain domains, in water between 6 to 9 eV and 20 to 200 eV, respectively (see the summery figure of cross sections in [3]), but due to dissociative electron attachment (DEA) these processes may extend to very low energies. The production of radiation due to de-excitation depends on excitation energy levels. For water molecule production of Lyman alpha radiation has a high cross section in the electron energy range from 50 to 200 eV.



Electron driven processes

Figure 1. Electron energy ranges that correspond to the relevant processes in collisions with atoms and molecules. Full colour corresponds to the specific case of water (data taken from Itikawa and Mason [3]) while the open bars are extended ranges that correspond to the majority of atomic species: red, elastic scattering; green, vibrational excitations; violet, attachment; blue, dissociation; orange, ionization; darkblue, Lyman alpha.

3. Atomic and Molecular Data Needed for Analysing Electron Scattering Processes Relevant to Comet 67P

Despite the evidence for electron induced processes in comets many of the discrete collision processes necessary to quantify such electron driven chemistry remain uncertain. In order to develop a predictive model of cometary coma and the comet's 'atmosphere' it is necessary to assemble a 'database' of relevant electron collision processes with the different atomic and molecular species observed (or indeed inferred) from Rosetta. As discussed above Rosetta's ROSINA, as well as Ptolemy and COSAC on the Philae lander, have revealed a rich chemical inventory that would require a large atomic and molecular physics database to model all possible processes. However, if a sensitivity analysis is performed the number of important species contributing to the model may be reduced to a minimum (more manageable) number of reactions. The dominant molecules are water, CO and CO₂. The presence of HCN as the source of CN radicals and ammonia as source of NH and NH₂ is widely accepted while the simple hydrocarbons C_2H_2 and C_2H_6 are assumed to be the source of C_2 , indeed C_2H_6 concentrations were unusually high in comet 67P [21]. Methane has been identified in many comets and is found in comet 67P. As discussed above several oxygen containing species were detected and apart from the ubiquitous water methanol CH₃OH may be an important primary compound. The primary source of sulphur compounds may be H₂S but 67P is depleted in all other sulphur bearing species $(CS_2, OCS \text{ and } SO_2)$ compared to other comets [22]. Thus, in developing an electron chemistry model of comet 67P it is necessary to have a good data base for electron interactions with H_2O , CO, CO₂, CH₄, C₂H₆, CH₃OH, NH₃, HCN and H₂S. The status of such a database will be discussed below but first it is necessary to understand the corollary for a 'good' database.

3.1. Databases

Many databases exist in order to assemble datasets and communicate them to different audiences. The NIST database collection (https://srdata.nist.gov/gateway/gateway?dblist=0) is one of the best known providing details of the structure, spectroscopy and fundamental parameters (ionization and dissociation energies) of many atoms and molecules. The need for large datasets has led to several communities investing in establishing data centres which assemble and maintain databases. For example, the fusion community has, for several decades, compiled databases in order to model plasmas in tokamak reactors and to provide data for diagnostic tools used in such plasmas. Another example of large collection of collisional data is the LXCaT database [23], which provides electron and ion scattering cross sections, swarm parameters (mobility, diffusion coefficient, etc.), reaction rates, energy distribution functions, etc. and other data required for modelling low temperature plasmas. Similarly, the astronomical community has needed large databases to interpret its observations, these include not only spectroscopic databases but also databases of chemical reaction rates (e.g., KIDA [24]) are necessary to understand the rich inventory of molecules that have been observed in the interstellar medium.

Many databases are simple collections of data but more recently the design and operation of databases has been refined. The development of IT tools has allowed data to be provided on-line, downloadable in a range of formats and allows new data to be added quickly, ensuring that the data is up to date. Previously data was reviewed and published in journal reviews which once published became gradually out of date until the next review, often a decade later (e.g., [25]). The opportunity to add new data quickly not only ensures that the latest data is adopted by the community but also reduces the likelihood of fragmentation amongst the community with different groups using different data sets in accord with their knowledge (or more commonly lack of knowledge) of the data available.

Simple assembly of data alone is not, however, the most effective form of databases. The 'user' community requires guidance as to what data to adopt. Users rarely have the necessary experience to select one dataset over another and therefore each may choose different sets, leading to systematic problems. For example, if different datasets are used in different models, cross comparison of such models is difficult and it may be hard to distinguish between the different physical and chemical

hypotheses in different models from the data used in the model to explore such effects. Therefore, databases should provide 'recommended data' which is the data that the expert community providing such data believes is the optimal data reflecting state of the art measurements or calculations. These values can be updated as new data becomes available. However, when changing recommended data, it is essential to ensure that it is still 'consistent' For example in presenting a comprehensive set of electron impact cross sections the individual cross sections (elastic and inelastic (including ionization excitation etc.) should, when summed, be consistent with the recommended data for total cross sections. Databases should also present data with stated estimates of uncertainties, particularly when presenting its own composite data from several different datasets.

3.2. VAMDC and BEAMDB Databases

The Virtual Atomic and Molecular Data Centre—(VAMDC) and Belgrade Electron Atom and Molecule Database—(BEAMDB) are two examples of new generation of databases. The VAMDC Consortium is a worldwide consortium which federates atomic and molecular databases through an e-science infrastructure to provide easy access to data from different databases via a single portal http://portal.vamdc.eu. About 90% of the inter-connected databases are focused on data that are used for the interpretation of astronomical spectra and for modelling in many fields of astrophysics and astrochemistry, although recently the VAMDC Consortium has connected databases from the radiation damage and the plasma communities which makes it suitable for medical and industrial applications [26]. While VAMDC does not itself select and analyse data it ensures data from its component databases are accessible in a single format and ascribe to general good practices as discussed above. The VAMDC Consortium includes new databases and services on a case by case basis during annual general scientific and technical meetings.

VAMDC provides its data in a XSAMS output. XSAMS is an XML representation of an atomic and molecular data model. The system allows for distributed querying of data via the VAMDC-TAP protocol, an implementation-agnostic standard, where data providers can build their models in their own fashion and map them to the VAMDC model via a dedicated dictionary [27].

The Belgrade Electron Atom and Molecule Database—(BEAMDB) [28] is an application, database and a VAMDC node which contains data for elementary processes of electron scattering by atoms and molecules. The database covers collisional data of electron interactions with atoms and molecules in the form of differential and integrated cross sections as well as energy loss spectra. The data is stored in a relational (MySQL) database, upon a static model specifically suited to this dataset but easily extendable. There have already been several migrations of the model, the latest of which is an extension to enable storing non-neutral molecules. Currently, there are 22 species stored in the database (11 atoms and 11 molecules), presented in 71 states, involved in 59 collision processes. The web interface (http://servo.aob.rs/emol) enables on-site querying of data via an AJAX-enabled web form. The application is implemented in Django, a Python web framework and hosted on an Apache web server at the Astronomical Observatory in Belgrade.

The BEAMDB is a collisional database where several types of collisions are included: Elastic, Electronic Excitation, (Total) Inelastic, Ionization, and Total Scattering as well as electron spectroscopic data such as Energy-loss Spectra and Threshold Photoelectron Spectra. Cross sections are of several different kinds: Differential, Integral, Total, Momentum Transfer, Viscosity. Specific data that are maintained in the BEAMDB are differential cross sections (DCS) for elastic scattering and excitation of atoms and molecules. These are 3D entries since DCS depend on both electron impact energy and scattering angle. This requires two X columns while the Y column is also associated with the column representing uncertainty of data points. An example of the XSAMS output of such kind of data for He excitation is shown in Figure 2.

Producing a plot of DCS data is not available at the current stage of database development, but such data can be visualized by using either VAMDC portal or alternatively the RADAM (RADiation DAMage) database portal (http://radamdb.mbnresearch.com/). The general structure of RADAM

databases covers electron/positron interactions, ionic and photonic interactions, multiscale radiation damage phenomena and radiobiological phenomena occurring at different time, spatial and energy scales in irradiated targets [29]. Examples of DCS surfaces for elastic electron scattering by helium atom and formamide (CH₃NO) molecule are shown in Figure 3a,b, respectively.

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Figure 2. XSAMS output table for differential cross section data for He electron excitation.



Figure 3. DCS surfaces as retrieved from RADAM database [29] for elastic electron scattering by: (a) He atom—data points are taken from ref. [30]; (b) Formamide molecule—data points are taken from ref. [31].

Initially BEAMDB was designed to maintain data of electron collisions with neutral species, atoms and molecules mainly in their ground state and exceptionally in excited state. However, by broadening the scope toward astrophysical applications, more specifically to the processes of comas that involve electrons, the database has been upgraded to include electron collisions with ions. The most recent data set included in BEAMDB is the one of electron ionisation cross sections of CN⁺ cations [32]. In the next few months BEAMDB will expand to include many more molecular systems including those needed to study the electron chemistry of comet 67P.

4. Electron Scattering Processes and Cross Sections—Data Needs

To understand and treat by models, processes in cometary plasmas is a very challenging task since many parameters need to be taken into account in order to cover the variety of comet types and their heterogeneity. One has to consider the changing of comet distances from the Sun and hence the level of irradiation and solar wind interactions both with the comet surface and cometary plasma environment. Nevertheless, models have been developed and set of processes reviewed, including data analysis used in such models. Particles from the solar wind, secondary electrons created in plasmas and photoelectrons, produce further events of excitation, ionization and dissociation with the consequence of enhanced chemical reactions and light emission. However, comets are composed of water, silicates and carbonaceous molecules (CO, CO₂ and hydrocarbons) [33]. Recently, modelling of plasma processes in cometary and planetary atmospheres has been performed by Campbell and Brunger [34] with an emphasis on the role of electron-impact excitation processes. They concluded in the case of comet Hale-Bopp that electron-impact could account for 40% of the fluorescence emissions of the fourth positive bands (A¹Π–X¹Σ⁺) of CO [35] and thus reducing calculated outgassing rates. Even more, their later paper [36] was focussed on electron initiated chemistry in atmospheres.

Reviews of cross section data and processes that cover electron scattering and excitations are numerous and they cover interactions with atoms [37,38], diatomic molecules [37,39], species in interstellar clouds [40] or concentrate on specific targets of triatomic molecules like water [3] or CO_2 and N_2O [40]. Anzai et al. [41] stressed that any recommended values of cross section data currently maintained in different databases might need to be updated due to the development of new experimental techniques and theoretical methods. The number of established benchmark cross sections is rather small.

The energy of the electrons available for electron interactions with atoms and molecules in the cometary coma 'atmosphere' is such that all electron scattering processes are relevant, thus a large amount of data is required if a model of electron induced processing is to be included in a simulation

model of comet 67P. In this section, the status of our knowledge of such relevant cross sections will be reviewed for the primary molecules defined in Section 3.

4.1. Elastic Electron Scattering and Cross Sections

Elastic scattering conserves the kinetic energy of the colliding particles. This means that quantum numbers that determine the energy are unchanged but other quantum numbers corresponding to degenerate states (e.g., helicity or spin flip) may change. In the case of many measurements not all states in the system are resolved due to the finite resolution of the electron beams used. In this case 'effective elastic cross sections' are determined which may be referred to as 'rotationally unresolved, vibrationally unresolved, electronically unresolved etc. Elastic scattering is important since, although there are no immediate changes in the target the range and hence spatial extent of the electrons is determined by such scattering.

Elastic cross sections are usually measured at specific energies and angles. These data are used to determine the 'total elastic cross section' by integrating over the entire angular range (4π). The total elastic cross section at given electron impact energy E_i is given by:

$$Q_{el}(E_i) = \int_0^{4\pi} \frac{d\sigma(k;\theta,\varphi)}{d\Omega} d\Omega,$$
(1)

where $\frac{d\sigma(k;\theta,\varphi)}{d\Omega}$ is the elastic differential cross section, (θ, φ) are the scattering angles and k is the wave-vector magnitude.

Elastic scattering is one of the best-studied electron collision processes and provides one of the best tests of developing theoretical calculations of electron-molecule scattering. The development of the magnetic angle changing method [42] to extend differential cross section measurements to the full range of scattering angles from 0° to 180° has significantly improved the accuracy of total elastic cross sections, particularly for molecules with dipole moments, where elastic scattering is strongly forward peaked.

Elastic scattering cross sections have been reported for all the primary molecules H_2O , CO, CO_2 , O_2 , CH_4 , C_2H_2 , CH_3OH , NH_3 , HCN and H_2S . Elastic scattering cross sections for water have been discussed in detail as part of a wider review of electron scattering from water by Itikawa and Mason [3]. An updated review has recommended the corrected data of Khakoo et al. ([43] and erratum) for low energy scattering and Munoz et al. [44] for higher energies (where experiment and theoretical evaluations merge). The benchmarking swarm paper by de Urquijo et al. [45] on cross sections for water reproduced measured transport data in water/helium mixtures and presented the integral cross sections that are entirely self-consistent with the available total cross sections as well as the swarm data over a large range of reduced electric field, E/N.

Cross sections for elastic scattering from methane and acetylene have recently been compiled and evaluated by Song et al. [46,47]. Compilations of data for other molecules are less recent and more fragmented and should be updated. Due to its toxicity, there are few measurements of the elastic scattering cross section from HCN and therefore there is more reliance on theoretical calculations (e.g., Sanz et al. [48]).

In elastic collisions electrons do not lose energy but change the direction of motion. This is important for models where the kinetics of all particles is taken into account. In more dense plasmas the elastic momentum transfer cross section, defined as integrated DCS with the weight of $(1 - \cos \theta)$ over all scattering angles, is a more relevant quantity. Differential cross sections, although being one of the basic properties that defines electron—atom/molecule interactions, are known with relatively low accuracy. In order to illustrate the current status of the agreement of DCS amongst different experiments and theories, the case of absolute cross sections for elastic electron scattering by argon atom is presented in Figure 4. It can be seen that although at first sight all values group around the averaged values, it should be noted that the data are plotted on a logarithmic scale and that there is almost an order of magnitude disagreement for particular data points.



Figure 4. DCS for elastic electron scattering by argon atom at 100 eV impact energy. The data symbols used correspond to: (red circles) , Rankovic et al. [49] experiment; (dark red full line), Rankovic et al. [49] theory; (magenta solid line), McEachran and Stauffer [50]; (olive dash dot line), Paikeday and Alexander [51]; (dotted line), Nahar and Wadehra [52]; (violet dash dot dot line), Fon et al. [53]; (cyan dash dot line), McCarthy et al. [54]; (dashed line), Joachain et al. [55]; (cyan up triangles), Milosavljević et al. [56]; (violet left triangle), Panajotović et al. [57]; (orange pentagons), Srivastava et al. [58]; (blue diamond), DuBois and Rudd [59]; (orange right triangles), Jansen et al. [60]; (green circles), Williams and Willis [61].

4.2. Electron Impact Ionisation Cross Sections

Most of the ions observed in the comet ion tail are the result of photoionisation of the primary ice species and since there are fewer high energy electrons, electron induced ionisation in comets is likely to be a minor process in total ion yields. However, the mass spectrometric analysis of compounds observed on Rosetta (e.g., using the Rosina instrument) rely upon knowing the fragmentation patterns of candidate molecules which, when compared with the instrument sensitivity, can be used to calculate the relative abundances of the detected molecules. Traditionally mass spectrometers operate with electron energies of 70 eV, close to but not at, the maximum of total ionisation cross sections. Branching ratios for fragments of electron impact ionisation are available in many databases (e.g., NIST Chemistry WebBook [62]) however, whilst these ratios are often known the cross sections are not presented. These cross sections may be derived if the total ionisation cross section is known. Total ionisation cross sections grave to an accuracy of <10% while semi-empirical calculations provide reliable cross sections (at least above 100 eV). Data on all of the primary molecules H₂O, CO, CO₂, CH₄, C₂H₂, CH₃OH, NH₃, HCN and H₂S exist with an accuracy sufficient for providing reliable data for determining their concentrations in the cometary coma and atmosphere. The recent review by
Tanaka et al. [63] covered this topic in detail employing scaled plane-wave Born models in order to provide comprehensive and absolute integral cross sections, first for ionization and then to optically allowed electronic-state excitation.

4.3. Anion Production

Anions have been observed in comets and may be formed by a variety of processes including radiative electron attachment, polar photodissociation, proton transfer and Dissociative Electron Attachment (DEA) where an incident electron is captured by the molecular target (AB) to form an excited state of the molecular negative ion AB⁻. This state, a Temporary Negative Ion (TNI), generally decays by ejecting the excess electron within a finite time (a process called autodetachment) but the molecular negative ion may also decay through dissociation leading to the formation of a stable negative ion B- and a neutral (often radical) fragment (A). DEA to the list molecules have been studied, identifying the fragment channels but there are few absolute cross sections. Node of VAMDC, the IDEADB maintained by University of Innsbruck, that serves data about dissociative electron attachment to molecules, lists more than 120 different fragments resulting from this process [64]. Anion data from comet 67P is still under evaluation but earlier studies from the Giotto spacecraft of comet 1P/Halley led to a combined chemical/hydrodynamic model for the coma of comet Halley to explore various anion production mechanisms and compute the abundances of atomic and molecular anions as a function of radius in the coma [65]. The dominant anion production mechanisms are found to be polar photodissociation of water and radiative electron attachment to carbon chains in the inner coma, followed by proton transfer from C_2H_2 and HCN to produce C_2H^- and CN^- , respectively. However, in the outer regions of the coma where electron temperatures reach 10^3 – 10^5 K, dissociative electron attachment may become a dominant process. Similar effects may be understood for comet 67P. DEA to water yields H^- and OH^- and O^- from CO and O_2 so there are many candidates for production of anions in comet 67P. DEA to all of the primary comet species (and most of the larger more complex species in Table 1) has been reported with DEA fragments recorded from near zero to the ionisation energy pathways. Nevertheless, very few absolute cross sections are available.

4.4. Electron Impact Excitation and Dissociation

Electron impact excitation and dissociative excitation of molecular systems is a critical process for a study of cometary coma and its tenuous atmosphere. As discussed above, OSIRIS and ALICE data from Rosetta shows the electron induced dissociation of water may be the source of the O I line at 135.6 nm while electron impact by CO and CO₂ yields C I lines at 156.1 and 165.7 nm. However, this hypothesis is handicapped by the dearth of data on electron impact excitation and electron induced neutral fragmentation for all molecules, not just those of immediate comet interest. This lack of experimental data can be attributed to difficulties in measuring neutral atoms and molecules. When an atom or molecule is in an excited state it may decay (fluoresce) with the emitted light being detected. Such experiments may identify some fragmentation or de-excitation pathways but the sensitivity of the optical detector and ability to 'capture' all of the emitted photons as well as the problem of cascades from higher lying states into the decaying state make measurements of absolute cross sections difficult. Furthermore, some excited states decay to 'dark' non-fluorescent or metastable states which makes them hard to detect. Although metastable fragments may be detected directly by surface ionization they will suffer from the same problems as photon detection i.e., cascade contributions may dominate [66,67]. Presently there are few experiments measuring electronically excited fragments by optical or metastable spectroscopy and more experiments are to be encouraged, building on the recent commissioning of electron induced fluorescence (EIF) experiment in Comenius University Bratislava. Figure 5 shows the H atom spectra recorded by electron impact of molecular hydrogen. This experiment is well equipped to study EIF of water, CO and CO₂ as required for cometary studies although, due to the low cross sections, data collection periods may be days or even weeks placing stringent conditions on the stability of the incident electron and gas beams.



Figure 5. Electron induced fluorescence of the H_2 molecule—Balmer lines and Fulcher α system. Measurements performed by Danko et al. [68].

Electron Energy loss spectroscopy monitors the energy of the incident electron post collision and may also be used to probe the direct excitation cross section of the parent molecule excited states but cannot provide data on the fragmentation patterns of that excited state as it decays. Furthermore, in most molecules the electronically excited states are both close together and overlap their ro-vibrational bands, making deconvolution very difficult if discrete electronic excitation cross sections are to be derived.

Finally, the production of neutral fragments in their ground state must be considered. Photon and electron induced dissociation produces many fragments in their ground state where ground state in this case includes fragments that are ro-vibrationally excited but still in the electronic ground state. This low internal energy precludes their detection by fluorescence since IR detection has not proven possible due to IR sources in the apparatus (e.g., electron filaments). Several alternative methods have been proposed to detect ground state neutral fragments including using a second electron beam to ionize the product, or use of surfaces to 'getter' the fragments. In the current context, there are only two experiments relevant to the modelling of electron dissociation of primary comet molecules—that of Harb et al. [69] measuring OH radical production from water and C and O from CO by Cosby et al. [70] using a fast beam method.

Further experimental studies on electron impact dissociation to neutral fragments will not only benefit the cometary community but the wider electron chemistry community with applications in many plasma systems, aeronomy and radiation chemistry. However, given the experimental difficulty much of the necessary data may be provided by theoretical calculations, which require more detailed exploration.

5. Possibility of Electron Induced Surface Chemistry

As discussed above (Section 2.3) many of the larger more complex molecules observed by COSAC and Ptolemy on the Philae lander may be made by addition reactions from simpler molecules. How

are such reactions induced? Photodissociation has been considered the primary process but electron induced chemistry within ices has been shown to be an efficient route to molecular synthesis and simple electron irradiation of primary ices has been shown to be produce most (all) of the larger molecular species. For example, Figure 6 shows the yield of formamide in an ice film composed of CO and NH₃ as a function of electron energy [71]. The ice was prepared with a mixing ratio of 1:8 and thickness corresponding to 12–18 monolayers and an electron exposure of 200 μ C/cm². Formamide is readily formed and the resonance like feature between 6 and 12 eV is characteristic of the synthesis by reactants prepared in a dissociative electron attachment process [71]. Similar experiments have shown that as many as 15 products can be formed by electron irradiation of pure methanol ices [72] including ethylene glycol and methyl formate whilst formamide HCONH₂ is formed in irradiation of binary mixtures of ammonia and methanol ice and the simplest amino acid glycine from irradiation of a methylamine and carbon dioxide ice [73]. Thus, electron induced synthesis of simple cometary ices may be a route to formation of several of the organic species observed in surface material from comet 67P.



Figure 6. The production of formamide present in mixed multilayer films of CO and NH₃. The resonance like feature (red line) between 6 and 12 eV is characteristic of the synthesis of formamide by reactants prepared in a DEA process, Bredehöft et al. [71].

The route by which molecular oxygen was formed as the fourth most abundant compound in the coma observed by Rosetta, is still subject to debate. However, laboratory experiments [74,75] have shown that radiolysis of water by both electrons and photons yields molecular oxygen but also copious amounts of hydrogen peroxide. Furthermore, comparative experiments between photon and electron irradiation show that electron induced yields are higher for the same energy. Whether this is due to penetration depth of electrons or that electrons open more dissociative pathways (through dipole or spin forbidden transitions) is unknown. The role of electron induced chemistry in comets, in ice covered planetary and lunar objects and in the rich chemistry of the interstellar medium is therefore an emerging topic of modern astronomy and one that has been encouraged by the results of the Rosetta mission.

6. Conclusions

In this paper, we have presented a review of recent results from the Rosetta mission to comet 67P/Churyumov-Gerasimenko. The role of electron induced processes has been highlighted with the emission of [O I] lines in the coma explained by the process of electron impact dissociation of water. The role of other electron processes e.g., in the production of the unexpectedly large amounts of molecular oxygen in the coma, is handicapped by lack of detailed knowledge of electron impact

cross sections We have reviewed the need for electron scattering data and discussed how such data should be collated and reviewed in electron scattering databases. The BEAMD database which is a part of a larger consortium of Virtual Atomic and Molecular Data Centre—VAMDC has been used as an example of modern generation of databases.

The importance of electron processes in comet 67P/Churyumov-Gerasimenko highlights the need for closer interactions and joint projects between the cometary and electron communities and this paper has identified some topics for joint research.

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Article



The Screening Characteristics of the Dense Astrophysical Plasmas: The Three-Component Systems

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Abstract: As the object of investigation, astrophysical fully ionized electron-ion plasma is chosen with positively charged ions of two different kinds, including the plasmas of higher non-ideality. The direct aim of this work is to develop, within the problem of finding the mean potential energy of the charged particle for such plasma, a new model, self-consistent method of describing the electrostatic screening. Within the presented method, such extremely significant phenomena as the electron-ion and ion-ion correlations are included in the used model. We wish to draw attention to the fact that the developed method is suitable for astrophysical applications. Here we keep in mind that in outer shells of stars, the physical conditions change from those that correspond to the rare, practically ideal plasma, to those that correspond to extremely dense non-ideal plasma.

Keywords: astrophysical plasmas; inner plasma electrostatic screening; different charged ions; stars

1. Introduction

Thematically, this work is the natural extension of the research on plasma's inner electrostatic screening, the results of which are presented in the papers [1–3]. In these papers, the single- and two-component systems are discussed with their properties in the region of higher non-ideality degree. This topic itself, the discussion and the search for more consistent models of screening and more realistic potentials in plasmas are still continuing and are very real (see [4–9]). The screening in astroplasma surroundings is a collective effect of many correlated particle interactions. It strongly affects the electronic structure, that is, the spectral properties of atoms and properties of their collision processes with respect to those for isolated systems [10,11]. In the last decade, a large number of theoretical as well as experimental investigations of plasma screening has been performed. For an example, it has been experimentally noted that the atomic spectral lines are redshifted in a high-power laser, producing dense plasmas as the result of these effects [12].

Here we consider, for the first time, systems of the next level of complexity, that is, three-component systems that contain free electrons and positively charged ions of two different kinds. Because of this, we recall that the conducted research had the following task: to investigate, within the problem of finding the mean potential energy of the charged particle in the plasma, whether the physical model of plasma's inner electrostatic screening, introduced in [13], is already exhausted by the Debye-Hückel (DH) method, as described in the same paper, or whether it still allows for the development of an alternative. As in the previous papers, here we keep in mind the electrostatic screening in fully ionized plasmas. Although the paper [13] was devoted to electrolytes, it had a profound trace in plasma physics and in adjacent disciplines [14–18]. Its influence is felt even today in various fields of physics,

such as ionospheric plasma physics, astrophysics, and laboratory plasma research [19–25]. Thus, in numerous papers, direct DH or DH-like methods are used, as well as their products such as the DH potential and DH radius (see [26–29]). This has all induced the interest for the possibility of going beyond the sphere of influence of [13] and for the development of the mentioned alternative method. Additionally, another stimulus exists for the development of alternative methods, connected with finding a characteristics length greater than the Debye radius ([30–35]; see also [3]).

We recall that the essential properties of the mentioned model are the following:

- The presence of an immobile probe particle, which represents one kind of charged particle in the real system (plasma, or electrolyte).
- The treatment of the considered components, which contain free charged particles of different kinds as ideal gases in states of thermodynamical equilibrium, without the assumption that all temperatures are equal.
- The treatment of the existing total electrostatic field in the considered system as an external field with respect to the considered ideal gas.
- Finally, among the properties of this model is usage, as its relevant mathematical apparatus, of equations, which describe the mean local electrostatic field and the conditions of conservation of thermodynamical equilibrium for the considered components. As in the previous papers, this model is treated here as the basic model.

The task formulated above itself has enforced a special referent role of the mentioned DH method, the predictions of which shall be compared with a possible new method that would arise as the result of the undertaken investigation. In accordance with this, the main aim of our previous research became the creation of a "self-consistent" method of describing the mentioned electrostatic screening mechanism, which is completely free of the DH method's disadvantages. We note that the definition of a self-consistent method implies that all the relevant characteristics are determined within this method itself and are expressed only through its basic parameters, that is, the particle densities, temperature, and so forth.

However, it has been shown that, except for the case of a single-component system (e.g., electron gas on a positively charged background), it was very difficult to finish the entire procedure of eliminating the disadvantages of the DH method in a self-consistent way. An analysis that was performed later convinced us that this result was not accidental, as the outer differences between the DH and the presented method were not practically significant, and the principal differences between these methods are of a conceptual nature.

Consequently, aside from finding the mean potential energy of the charged particles in the plasmas, the direct objective of this research became the development of a self-consistent method of describing the electrostatic screening in the considered three-component system, for which the relevant additional conditions (equivalent to the conditions of the conservation of particle numbers in finite systems) are included from the beginning.

In order to show the differences between the results of applying and neglecting the relevant additional conditions, we recall the behaviour of electron density in two cases of the electron-ion plasmas with the probe particle whose charge is equal to that of an ion, which was considered in [2] and is illustrated here in Figure 1a,b. The first of these illustrates the application of the method developed in [2], which makes sure that the area of higher electron density is followed by an area of its lowering. This is in accordance with the role of the probe-particle approximation: the situation in the vicinity of the probe particle could be considered as a reflection of the real situation in the vicinity of any ion in the considered plasma, and it should enable usage of the results illustrated by the figure to the case of the real plasma, as these do not influence the mean electron density. The systems with similar behaviour in the electron density are treated here as the "closed" systems. In Figure 1b, the behaviour of the DH electron density is shown in electron–proton plasma with the probe particle whose charge is equal to that of the proton. From this figure, a monotonous increasing of the electron density can be

seen, with a decrease in the distance from the probe particle from infinity to zero. In the considered case, such behaviour causes the creation of an excess of 1/2 electron and 1/2 proton in the vicinity of the probe particle. This phenomenon, which is unacceptable from the point of view of a method that includes additional conditions, is discussed in detail in Section 6.



Figure 1. (a) The behavior of the electron density in the case of the electron–ion plasma with the probe particle whose charge is equal to that of the ions; (b) the behavior of the Debye-Hückel (DH) electron in the case of the electron-proton plasma with the probe particle whose charge is equal to that of the protons.

This work is dedicated to plasmas that are treated as fully ionized, including the plasmas of higher non-ideality. The region of electron densities from 10^{16} to 10^{20} cm⁻³ and temperatures from 1×10^4 to 3×10^4 K are studied. The developed theory is also applicable to a wider area of plasma parameters. The order of its exposing mostly follows that which is presented in our previous papers [1–3].

2. Theory Assumptions

2.1. The Initial System and Basic Characteristic

A stationary homogeneous and isotropic system S_{in} is taken here as the initial model of some real physical objects, suitable for applications of the results of this research. It is assumed that S_{in} is constituted by a mixture of a gas of free electrons and two gases of free ions of different kinds with positive charges Z_1e and Z_2e , where $Z_{1,2} = 1, 2, 3, ...$, and e is the modulus of the electron charge. The electron charge -e is denoted also by Z_ee , where $Z_e = -1$. We consider that these gases are in equilibrium states with mean densities of N_1 and N_2 and temperatures of $T_1 = T_2 = T_i$ for the ions, and a mean density of N_e and temperature of $T_e \ge T_i$ for the electrons. All the particles are treated as point-like, non-relativistic objects and their spins are taken into account only as factors that influence the chemical potentials of the considered gases. Satisfying the condition

$$Z_1 e \times N_1 + Z_2 e \times N_2 - e \times N_e = 0 \tag{1}$$

is assumed, which provides local quasi-neutrality of the system S_{in} . We emphasize that the case $Z_1 = Z_2$ is also considered here (see Section 6.1), as it reflects the existence of some real systems with two physically different kinds of ions with the same charge, for example, H⁺ and D⁺, or H⁺ and He⁺, and so forth.

In the subsequent considerations, several characteristic lengths are used that are connected with the parameters $N_{1,2,e}$ and $Z_{1,2}$, namely, the Wigner–Seit's (WS) radii and the "ion self-spheres" radii (see [3]), denoted here by $r_{1,2,e}$ and $r_{s;1,2}$, respectively. These are defined by the relations

$$\frac{4\pi}{3} \times r_{1,2,e}^3 = \frac{1}{N_{1,2,e}}, \quad \frac{4\pi}{3} \times r_{s;1,2}^3 = \frac{Z_{1,2}}{N_e}, \quad r_{s;e} \equiv r_e \tag{2}$$

From here, it follows that Equation (1) can be presented in the form

$$p_1 + p_2 = 1, \qquad p_{1,2} \equiv \frac{N_{1,2} \times Z_{1,2}}{N_e} = N_{1,2} \times \frac{4\pi}{3} r_{s;1,2}^3,$$
 (3)

where the parameters p_1 and p_2 describe the primary distribution of the space between the self-spheres of all ions of the first kind and all ions of the second kind.

2.2. The System Properties and Conditions

In accordance with the basic model and the composition of the system S_{in} , the electrostatic screening of the charged particles is modeled here in three corresponding auxiliary systems. It is assumed that each of these contains the following: the electron component, two ion components with the same charges Z_1e and Z_2e , and one immobile probe particle with the charge Z_pe , which is fixed at the origin of the used reference frame (the point *O*).

As in [1,2], only such cases are studied here for which the probe particle can represent one of the charged particles of the system S_{in} , for example, when $Z_p = Z_1$, Z_2 and Z_e . Two ion cases are denoted below with (*i*1) and (*i*2), and the electron case-with (*e*), while the corresponding auxiliary systems are denoted with $S_a^{(1)}$, $S_a^{(2)}$ and $S_a^{(e)}$, respectively.

All systems $S_a^{(1,2,e)}$ are treated below as isotropic and are characterized by the corresponding mean local ion and electron densities: $n_1^{(1,2,e)}(r)$, $n_2^{(1,2,e)}(r)$ and $n_e^{(1,2,e)}(r)$, which retain the properties of the corresponding components in the system S_{in} and satisfy the boundary conditions:

$$\lim_{r \to \infty} n_1^{(1,2,e)}(r) = N_1, \quad \lim_{r \to \infty} n_2^{(1,2,e)}(r) = N_2, \quad \lim_{r \to \infty} n_e^{(1,2,e)}(r) = N_e \tag{4}$$

where $r = \vec{r}$, and \vec{r} is the radius vector of the observed point. Their other necessary characteristics are the mean local charge density $\rho^{(1,2,e)}(r)$ defined by the relation

$$\rho^{(1,2,e)}(r) = Z_1 e \times n_1^{(1,2,e)}(r) + Z_2 e \times n_2^{(1,2,e)} - e \times n_e^{(1,2,e)}(r)$$
(5)

and the mean local electrostatic potential $\Phi^{(1,2,e)}(r)$, which is treated as the potential of the external electrostatic field. We take into account the fact that $\Phi^{(1,2,e)}(r)$ and $\rho^{(1,2,e)}(r)$ have to satisfy Poisson's equation:

$$\nabla^2 \Phi^{(1,2,e)} = -4\pi \left[Z_{1,2,e} e \times \delta(\vec{r}) + \rho^{(1,2,e)}(r) \right]$$
(6)

where $\delta(\vec{r})$ is the three-dimensional delta function [14], and $0 \leq r \leq \infty$. Satisfying the boundary conditions

$$\lim_{r \to \infty} \Phi^{(1,2,e)}(r) = 0, \quad \left| \varphi^{(1,2,e)} \right| < \infty; \varphi^{(1,2,e)} \equiv \lim_{r \to 0} [\Phi^{(1,2,e)}(r) - \frac{Z_{1,2,e}e}{r}]$$
(7)

is assumed, which guaranties a physical sense of the mentioned electrostatic potential and connection with the system S_{in} and is compatible with the electro-neutrality condition of the auxiliary systems. Because $\varphi^{(1,2,e)}$ is the mean electrostatic potential at the point O, the quantity

$$U^{(1,2,e)} = Z_{1,2,e} e \times \varphi^{(1,2,e)} \tag{8}$$

is the mean potential energy of the probe particle and is simply the searched mean potential energy of the probe particle. In the usual way, $U^{(1,2)}$ and $U^{(e)}$ are treated as approximations to the mean potential energies of the ion and electron in the initial system S_{in} .

In accordance with the basic model the electron, all ion components of all auxiliary systems are treated as ideal gases. Therefore, we encompass the characteristics of the auxiliary systems by chemical potentials $\mu_{1,2}(n_{1,2}(r), T_i)$ and $\mu_e(n_e^{(e)}(r), T_e)$ of the corresponding ideal ion and electron gases, which can depend on the corresponding particle spins and on their boundary values, that is, $\mu_{1,2}(N_{1,2}^{(1,2)}, T_i) = \lim_{r \to \infty} \mu_{1,2}(n_{1,2}(r), T_i) = \lim_{r \to \infty} \mu_{1,2}(n_{1,2}^{(2,1)}(r), T_i)$ and $\mu_e(N_e, T_e) = \lim_{r \to \infty} \mu_e(n_e^{(e)}(r), T_e)$.

2.3. The System Equations

It can be shown that on the basis of the procedure that was developed and described in detail in [1,2] for the case of a system that is electro-neutral as a whole, it is possible to switch from Poisson's equation to the equation for the potential $\Phi^{(1,2,c)}(r)$, which is more suitable for further consideration. This equation is given by

$$\Phi^{(1,2,e)}(r) = -4\pi \int_{r}^{\infty} \rho^{(1,2,e)}(r)(r') \left(\frac{1}{r} - \frac{1}{r'}\right) r'^2 dr'$$
(9)

and is taken here in such a form.

In order to find other necessary equations, we consider the conditions of conservation of thermodynamical equilibrium (conservation of the electro-chemical potential) for those components that are represented by the corresponding probe particles, namely,

$$\mu_{\delta}(n_{\delta}^{(\delta)}(r), T_{\delta}) + Z_{\delta}e \times \Phi^{(\delta)}(r) = \mu_{\delta}(n_{\delta}^{(\delta)}(r_{st}), T_{\delta}) + Z_{\delta}e \times \Phi^{(\delta)}(r_{st}), \quad \delta = 1, 2, e$$
(10)

where, in accordance with the basic model, $\Phi^{(1,2,e)}(r)$ is treated as the potential of the external electrostatic field, and r_{st} is the distance from the point *O* of the chosen fixed (starting) point:

 $0 < r_{st} \leq \infty$. From here, by means of the usual linearization procedure, the necessary equations for the particle densities $n_{\delta}^{(\delta)}(r)$ are obtained in the form

$$n_{\delta}^{(\delta)}(r) - n_{\delta}^{(\delta)}(r_{st}) = -\frac{Z_{\delta}e}{\partial\mu_{\delta}/\partial N_{\delta}} \times \left[\Phi^{(\delta)}(r) - \Phi^{(\delta)}(r_{st})\right], \quad \frac{\partial\mu_{\delta}}{\partial N_{\delta}} \equiv \left[\frac{\partial\mu_{\delta}(n, T_{\delta})}{\partial n}\right]_{n=N_{\delta}} \tag{11}$$

which is applicable under the condition

$$\frac{|n_{\delta}^{(\delta)}(r) - N_{\delta}|}{N_{\delta}} \ll 1 \tag{12}$$

Here we use the fact that such equations can be applied not only to the classical cases, but also to the quantum-mechanical cases, including the case of ultra-degenerated electron gas [15,17].

2.4. The Additional Conditions

In [1,2], the conditions were already introduced for the component that is represented by the probe particle and in which the charge of the probe particle appears. In the considered three-component case, these conditions are given by

$$\int_{0}^{\infty} \left[N_{1,2,e} - n_{1,2,e}^{(1,2,e)}(r) \right] \times 4\pi r^2 dr = \int_{0}^{\infty} \left[1 - \frac{n_{1,2,e}^{(1,2,e)}(x \times r_{1,2,e})}{N_{1,2,e}} \right] \times 3x^2 dx = 1$$
(13)

where $x \equiv r/r_{1,2,e}$. This means that the ratio in the expression inside the square brackets is of the order of magnitude of 1. This equation is especially important, as it provides the continuity of the model. In order to show this fact, it is enough to consider the situation for which the charge density $n_{1,2}^{(2,1)}$ is negligible and the considered three systems for physical reasons can be treated as a two-component system. Then, as an approximation, we can replace the electron component by the negatively charged nonstructural background and return to a one-component system. It is important that the single-component case is now used for mathematical modeling of the plasma internal electrostatic screening (Iosilevskiy 2011, private communication). We note that in the single-component case (which was not considered in [13]), this first condition can be used instead of the electro-neutrality condition.

In [2], the additional conditions for the electron density were introduced. Here the corresponding conditions are given by the equations

$$\int_{0}^{\infty} \left[N_{1,2} - n_{1,2}^{(e)}(r) \right] \times 4\pi r^2 dr = \int_{0}^{\infty} \left[N_e - n_e^{(1,2)}(r) \right] \times 4\pi r^2 dr = 0$$
(14)

The additional conditions for the ion components $n_1^{(2)}(r)$ and $n_2^{(1)}(r)$ are taken into consideration analogously to the electron components, as there are no principal differences between them. The corresponding relations are given by

$$\int_{0}^{\infty} \left[N_1 - n_1^{(2)}(r) \right] \times 4\pi r^2 dr = \int_{0}^{\infty} \left[N_2 - n_2^{(1)}(r) \right] \times 4\pi r^2 dr = 0$$
(15)

We note that in this work, the considered physical systems, as well as other physical systems that are described by means of additional conditions, are treated as systems of the closed type (see also a detailed version [36]).

As it is known, in our investigation, we take care that our results are compared with the results of the DH method, which provides electro-neutrality of the considered system as a whole.

In principle, this would justify introducing into consideration the corresponding electro-neutrality condition, namely,

$$Z_{1,2,e}e + \int_{0}^{\infty} \rho^{(1,2,e)}(r) \times 4\pi r^2 dr = 0$$
(16)

However, the fact is used that simultaneous satisfaction of the conditions of Equations (4), (14) and (40) automatically provides satisfaction of this condition, and therefore this condition is not used within this work.

3. Ion Cases: Complete Expressions

3.1. The Ion Densities

As a result of their importance, the ion densities are presented separately for the case (i1) in the form (1)

$$n_{1}^{(1)}(r) = N_{1} \times \begin{cases} 0, & 0 < r \le r_{0;1}^{(1)} \\ 1 - A_{1} - B_{1}d_{1}r_{b;1} \times \frac{F_{1}(r)}{r}, & r_{0;1}^{(1)} < r \le r_{b;1} \\ 1 - C_{1}r_{b;1} \times \frac{e^{-\kappa_{as;1}(r-r_{b;1})}}{r}, & r_{b;1} < r < \infty \end{cases}$$
(17)

$$n_{2}^{(1)}(r) = N_{2} \times \begin{cases} 0, & 0 < r \le r_{0,2}^{(1)} \\ 1 + \frac{N_{1}Z_{1}}{N_{2}Z_{2}} \times A_{1} - \frac{N_{1}Z_{1}}{N_{2}Z_{2}} \times B_{1}d_{2}r_{b;1} \times \frac{F_{1}(r)}{r}, & r_{0;2}^{(1)} < r \le r_{b;1} \\ 1 + \frac{N_{1}Z_{1}}{N_{2}Z_{2}} \times C_{1}\alpha_{i}r_{b;1} \times \frac{e^{-\kappa_{a;1}(r-r_{b;1})}}{r}, & r_{b;1} < r < \infty \end{cases}$$
(18)

and separately for the case (i2) in the similar form

$$n_{2}^{(2)}(r) = N_{2} \times \begin{cases} 0, & 0 < r \le r_{0;2}^{(2)} \\ 1 - A_{2} - B_{2}d_{2}r_{b;2} \times \frac{F_{2}(r)}{r}, & r_{0;2}^{(2)} < r \le r_{b;2} \\ 1 - C_{2}r_{b;2} \times \frac{e^{-\kappa_{as;2}(r-r_{b;2})}}{r}, & r_{b;2} < r < \infty \end{cases}$$
(19)

$$n_{1}^{(2)}(r) = N_{1} \times \begin{cases} 0, & 0 < r \le r_{0;1}^{(2)} \\ 1 + \frac{N_{2}Z_{2}}{N_{1}Z_{1}} \times A_{2} - \frac{N_{2}Z_{2}}{N_{1}Z_{1}} \times B_{2}d_{1}r_{b;2} \times \frac{F_{2}(r)}{r}, & r_{0;1}^{(2)} < r \le r_{b;2} \\ 1 + \frac{N_{2}Z_{2}}{N_{1}Z_{1}} \times C_{2}\alpha_{i}r_{b;2} \times \frac{e^{-\kappa_{as;2}(r-r_{b;2})}}{r}, & r_{b;2} < r < \infty \end{cases}$$
(20)

where the functions $F_{1,2}(r)$ and the coefficients $f_{1,2}$, $A_{1,2}$, $B_{1,2}$, $C_{1,2}$ and $d_{1,2}$ are given by the relations

$$F_{1,2}(r) \equiv e^{\kappa_i(r_{b;1,2}-r)} - f_{1,2} \times e^{-\kappa_i(r_{b;1,2}-r)}$$

$$f_{1,2} = \frac{(1 + \kappa_{as;1,2}r_{b;1,2}) \times \kappa_i^2 - (1 + \kappa_i r_{b;1,2}) \times \kappa_{as;1,2}^2}{(1 + \kappa_{as;1,2}r_{b;1,2}) \times \kappa_i^2 - (1 - \kappa_i r_{b;1,2}) \times \kappa_{as;1,2}^2}$$
(21)

$$A_{1,2} = [1 - d_{1,2}(1 - \alpha_i)] \times C_{1,2}, \qquad B_{1,2} = \frac{1 - \alpha_i}{1 - f_{1,2}} \times C_{1,2}$$
(22)

$$C_{1,2} = \left[1 - d_{1,2}(1 - \alpha_i) \left(1 - \frac{r_{b;1,2}}{1 - f_{1,2}} \times \frac{F_{1,2}(r_{0;1,2}^{(1,2)})}{r_{0;1,2}^{(1,2)}} \right) \right]^{-1}, \quad d_{1,2} = \frac{\kappa_{0;1,2}^2}{\kappa_{0;1}^2 + \kappa_{0;2}^2}$$
(23)

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and the screening constants

$$\kappa_{as;1,2} = \kappa_{0;1,2} \times \left[(1 - \alpha_{e;1,2}) \times (1 - \alpha_i) \right]^{1/2}, \quad \kappa_{0;1,2} = \left[\frac{4\pi (Z_{1,2}e)^2}{\partial \mu_{1,2} / \partial N_{1,2}} \right]^{1/2}$$
(24)

$$\kappa_i = [(\kappa_{0;1}^2 + \kappa_{0;2}^2) \times (1 - \alpha_{e;1,2})]^{1/2}$$
(25)

3.2. The Electron Densities

The complete expressions for the electron densities are presented here in the form

$$n_{e}^{(1,2)}(r) = \alpha_{e;1,2} \times [Z_1 \times n_1^{(1,2)}(r) + Z_2 \times n_2^{(1,2)}(r)] + \begin{cases} n_{s,e}^{(1,2)}(r), & 0 < r \le l_{s;1,2} \\ N_e \times (1 - \alpha_{e;1,2}), & l_{s;1,2} < r < \infty \end{cases}$$
(26)

where the ion densities $n_1^{(1,2)}(r)$ and $n_2^{(1,2)}(r)$ are given by Equations (17)–(19). The number $n_{s;e}^{(1,2)}(r)$, in accordance with the above, is given by the relations

$$n_{s;e}^{(1,2)}(r) = N_e \times l_{s;1,2} \frac{a_{1,2} \times e^{-\kappa_{0;e}r} + b_{1,2} \times e^{\kappa_{0;e}r}}{r}, \qquad 0 < r < l_{s;1,2}$$
(27)

$$a_{1,2} = \frac{1 - \alpha_{e;1,2} - \frac{1}{3}x_{l;1,2}^2 \times e^{x_{l;1,2}}}{e^{-x_{l;1,2}} - e^{x_{l;1,2}}}, \qquad b_{1,2} = -\frac{1 - \alpha_{e;1,2} - \frac{1}{3}x_{l;1,2}^2 \times e^{-x_{l;1,2}}}{e^{-x_{l;1,2}} - e^{x_{l;1,2}}}$$
(28)

$$x_{l;1,2} = \kappa_{0;e} \times l_{s;1,2}, \qquad \kappa_{0;e} = \left[\frac{4\pi e^2}{\partial \mu_e / \partial N_e}\right]^{1/2}$$
 (29)

We note that these parameters, as well as $r_{b;1,2}$, $r_{0;1}^{(1,2)}$, $r_{0;2}^{(1,2)}$ and $r_{0;1}^{(1,2)}$, are determined as described in Section 5.

4. Complete Expressions for the Electron and Ion Densities: The Case (e)

Here we can repeat the procedures from [2] verbatim. We obtain the expression for the electron density:

$$n_{e}^{(e)}(r) = \begin{cases} N_{e} - N_{e}r_{0;e} \times \exp(\kappa_{e}r_{0;e}) \times \frac{\exp(-\kappa_{e}r)}{r}, & r_{0;e} < r < \infty \\ 0, & 0 < r \le r_{0;e} \end{cases}$$
(30)

which determines $n_e^{(e)}(r)$ in the whole region $0 < r < \infty$. The obtained expressions are given here by the relations

$$n_{1,2}^{(e)}(r) = \frac{\alpha_{e;1,2}p_{1,2}}{Z_{1,2}} \times n_e^{(e)}(r) + \begin{cases} N_{1,2}(1 - \alpha_{e;1,2}), & l_{e;1,2} < r < \infty\\ n_{e;1,2}^{(e)}(r), & 0 < r \le l_{e;1,2} \end{cases}$$
(31)

$$n_{s;1,2}^{(e)}(r) = N_{1,2}r_e \frac{a_{1,2} \times e^{-\frac{x_{l;1,2} \times r}{l_{c;1,2}}} + b_{1,2} \times e^{\frac{x_{l;1,2} \times r}{l_{c;1,2}}}}{r}, \qquad 0 < r \le l_{e;1,2}$$
(32)

$$l_{e;1,2} = r_e \times \frac{l_{s;1,2}}{r_{s;1,2}} \tag{33}$$

where $a_{1,2}$ and $b_{1,2}$ are given by Equation (28). We note that the ratios $l_{s;1,2}/r_{s;1,2}$ and the characteristic length $r_{0;e}$ are determined as described in the next section.

5. Determination of the Parameters

The parameters $\alpha_{e;1,2}$ and $l_{s;1,2}$ are determined separately for the cases $Z_1 = Z_2 = Z_i$ and $Z_1 \neq Z_2$; namely, within the used procedure, the first case, where $r_{s;1} = r_{s;2} \equiv r_{s;i}$, is equivalent (from the point of view of the determination of $\alpha_{e;1,2}$ to the case of the two-component plasma with the same Z_i , N_e and T_e . Consequently, in the case for which $Z_{1,2} = Z_i$, the following relations are valid:

$$l_{s;1,2} = r_{s;i}, \qquad \alpha_{e;1,2} = \alpha(x_{s;i}), \quad x_{s;i} \equiv \kappa_{0;e} \times r_{s;i}$$
 (34)

where in accordance with [2], $\alpha(x)$ is defined by

$$\alpha(x) = 1 - \frac{\frac{2}{3}x^3}{(1+x) \times e^{-x} - (1-x) \times e^x}$$
(35)

In the case for which $Z_1 \neq Z_2$, the parameters $\alpha_{e;1,2}$ are given by the relation

$$\alpha_{e;1,2} \cong \alpha(x_s = x_{s;1}) \times p_1 + \alpha(x_s = x_{s;2}) \times p_2, \quad |l_{s:1,2}/r_{s;1,2} - 1| << 1$$
(36)

and are established by direct calculations where p_1 and p_2 are given by Equations (2) and (3).

It is important that the electron–ion correlation coefficient $\alpha_{e;1,2}$ and the characteristic lengths $l_{s;1,2}$ are determined, as is described below, independently of all other parameters; namely, this is why the existing conditions are sufficient for the determination of the characteristic lengths $r_{0;1,2,e}^{(1,2,e)}$, $r_{0;2,1}^{(1,2)}$ and $r_{b;1,2}$, and the ion–ion correlation coefficient α_i .

The very important parameters $r_{b;1,2}$, that is, the distances from the point *O* at which the manner of describing the ion densities changes, are determined from

$$\int_{0}^{\infty} \left[N_1 - n_1^{(2)}(r) \right] \times 4\pi r^2 dr = \int_{0}^{\infty} \left[N_2 - n_2^{(1)}(r) \right] \times 4\pi r^2 dr = 0$$
(37)

through a procedure for which it is taken that

$$r_{b;1,2} = r_{s;1,2} \times (1 + \eta_{1,2}), \quad 0 < \eta_{1,2} \le \eta_{max;1,2}, \quad \eta_{max;1,2} \gg 1$$
(38)

where $\eta_{1,2}$ are new parameters, which are used in the calculations in such a way that they vary with the small steps $\Delta \eta_{1,2} = 1/K_{1,2}$ where $K_{1,2} \gg 1$. As the results of this procedure, we obtain the values of the parameters $r_{b;1,2}$, that is, the main considered characteristic length, which corresponds to the current value of the ion–ion correlation coefficient α_i . The final value of this coefficient itself is determined through a procedure that implies scanning α_i with a very small step in the interval from 0 to 1 and examining at each step whether the equation

$$r_{0:1}^{(2)} - r_{0:2}^{(1)} = 0 (39)$$

is satisfied, which provides the physical meaning of the obtained solutions. The whole procedure ends when the equation is satisfied.

The parameter $r_{0:e}$ is determined from

$$\int_{0}^{\infty} \left[N_{1,2,e} - n_{1,2,e}^{(1,2,e)}(r) \right] \times 4\pi r^2 dr = \int_{0}^{\infty} \left[1 - \frac{n_{1,2,e}^{(1,2,e)}(t \times r_{1,2,e})}{N_{1,2,e}} \right] \times 3t^2 dt = 1$$
(40)

as in [1,2], using $t = r/r_{1,2,e}$ and Equation (2). It can be presented in two equivalent forms:

$$r_{0;e} = \frac{(1+x^3)^{\frac{1}{3}} - 1}{x} \times r_e \equiv \gamma_{s;e}(x) \times r_e, \qquad r_{0;e} = [(1+x^3)^{\frac{1}{3}} - 1] \times r_{\kappa;e} \equiv \gamma_{\kappa;e}(x) \times r_{\kappa;e}$$
(41)

where $x = \kappa_e r_e$, $r_{\kappa;e} \equiv 1/\kappa_e$, and the coefficients $\gamma_{s;e}(x)$ and $\gamma_{\kappa;e}(x)$ are connected with the electron non-ideality parameters $\Gamma_e = e^2/(kT_e r_e)$ and $\gamma_e = e^2/(kT_e r_{\kappa;e})$ as described in [3].

Finally, we note that the partial electron and ion densities $n_{s;e}^{(1,2)}$ and $n_{s;1,2'}^{(e)}$ because of the structure of the equation for the coefficients $a_{1,2}$ and $b_{1,2}$ (Equation (28)), can be determined by Equations (27), (29) and (33) in both the $Z_{1,2} = Z_i$ and $Z_1 \neq Z_2$ cases. Therefore, it is necessary to take the corresponding values of $\alpha_{e;1,2}$ and $l_{s;1,2}$ only in these expressions, for example, $l_{s;1,2} = r_{s;i}$ if $Z_{1,2} = Z_i$.

The behaviour of the characteristic length $l_{s;1,2}$, the electron–ion correlation coefficients $\alpha_{e,1,2}$, the parameters $r_{b;1,2}$ and the ion–ion correlation coefficients α_i is shown in Tables 1 and 2. These tables cover the regions of N_e from 10^{16} to 10^{20} cm⁻³ for $T = 3 \times 10^4$ K. These tables show that the values of all the parameters are within the expected boundaries. Additionally, one may note that particularly for $l_{s;1,2} \approx r_{s;1,2}$, the electron–ion correlation coefficient $l_{e;1,2} \approx 1$, $r_{b;1,2} \sim r_{s;1,2}$ and the ion–ion correlation coefficient $\alpha_i \approx 1$, a significant increase in the correlation coefficient values $\alpha_{e,1,2}$ and α_i is registered in the region of extremely high electron density ($N_e \approx 10^{19}$ cm⁻³).

Table 1. The characteristic length $l_{s;1,2}$ (in 10^{-7} cm), the non dimensional electron–ion correlation coefficients $\alpha_{e,1,2}$, the main characteristic length $r_{b;1,2}$ (in 10^{-7} cm), the non-dimensional ion–ion correlation coefficients α_i and the potential energies $U^{(1)}$ (in eV) and $U^{(2)}$ (in eV) for the cases of $Z_1 = 1$ and $Z_2 = 2$ at $T = 3 \times 10^4$ K in the region of electron densities 10^{16} cm⁻³ $\leq N_e \leq 10^{20}$ cm⁻³. The densities $N_{1,2}$ are in 10^{17} cm⁻³.

N_1	N_2	$l_{s;1}$	$l_{s;2}$	$\alpha_{e,1}$	$\alpha_{e,2}$	$r_{b;1}$	$r_{b;2}$	α _i	$U^{(1)}$	<i>U</i> ⁽²⁾
0.1	0.45	14.53	16.71	0.01	0.02	22.96	73.37	0.10	-1.09	-1.19
0.2	0.4	14.42	16.58	0.01	0.02	23.07	55.56	0.09	-2.79	-2.84
0.3	0.35	14.30	16.45	0.01	0.02	24.60	47.87	0.09	-2.39	-2.43
0.4	0.3	14.19	16.31	0.01	0.02	26.53	42.90	0.09	-2.14	-2.17
0.5	0.25	14.06	16.17	0.01	0.02	30.51	41.40	0.10	-2.07	-2.09
0.6	0.2	13.93	16.02	0.01	0.02	30.23	34.29	0.08	-1.71	-1.72
0.7	0.15	13.80	15.87	0.01	0.02	32.15	29.99	0.07	-1.60	-1.61
0.8	0.1	13.66	15.71	0.01	0.02	41.26	30.32	0.08	-2.07	-2.07
0.9	0.05	13.52	15.54	0.01	0.02	55.42	28.75	0.08	-2.83	-2.78
1	4.5	6.74	7.76	0.03	0.05	10.24	29.40	0.18	-1.35	-4.77
2	4	6.69	7.70	0.03	0.05	10.10	22.79	0.16	-3.45	-3.62
3	3.5	6.63	7.64	0.03	0.05	10.41	19.40	0.15	-2.89	-3.00
4	3	6.58	7.58	0.03	0.05	11.18	17.57	0.15	-2.64	-2.71
5	2.5	6.52	7.51	0.03	0.05	12.59	16.75	0.16	-2.53	-2.58
6	2	6.46	7.44	0.03	0.05	14.41	16.22	0.17	-2.46	-2.49
7	1.5	6.40	7.37	0.03	0.05	10.31	9.66	0.07	-3.32	-3.25
8	1	6.34	7.30	0.03	0.04	16.16	12.04	0.12	-2.47	-2.44
9	0.5	6.27	7.22	0.03	0.04	22.20	11.92	0.13	-3.48	-3.36
10	45	3.12	3.60	0.06	0.09	4.93	12.25	0.34	-5.52	-5.56
20	40	3.10	3.58	0.06	0.09	4.71	9.62	0.29	-4.04	-4.25
30	35	3.07	3.55	0.06	0.09	4.88	8.48	0.28	-3.49	-3.75
40	30	3.05	3.52	0.06	0.09	5.12	7.67	0.27	-3.25	-3.42
50	25	3.02	3.49	0.06	0.09	5.56	7.15	0.27	-3.07	-3.20
60	20	3.00	3.46	0.06	0.09	8.12	8.92	0.43	-3.99	-4.00
70	15	2.97	3.43	0.06	0.09	3.86	3.60	0.09	-3.44	-3.38
80	10	2.94	3.39	0.06	0.09	5.88	4.45	0.15	-2.55	-6.03
90	5	2.91	3.36	0.06	0.09	8.35	4.67	0.18	-3.77	-3.37
100	450	1.44	1.67	0.12	0.18	3.01	5.85	0.70	-6.89	-6.16
200	400	1.43	1.66	0.12	0.18	3.79	5.93	0.81	-7.36	-6.44
300	350	1.42	1.65	0.12	0.18	4.49	6.13	0.87	-2.11	-7.01
400	300	1.41	1.64	0.12	0.18	4.18	5.37	0.81	-6.46	-5.75
500	250	1.40	1.62	0.12	0.18	3.79	4.51	0.71	-5.00	-4.66
600	200	1.39	1.61	0.12	0.18	4.08	4.38	0.71	-4.88	-4.61
700	150	1.37	1.60	0.12	0.18	1.83	1.72	0.17	-3.91	-3.10
800	100	1.36	1.58	0.12	0.18	2.06	1.60	0.16	-4.86	-3.86
900	50	1.35	1.57	0.12	0.18	3.00	1.75	0.21	-3.20	-3.06

N_1	N_2	$l_{s;1}$	$l_{s;2}$	$\alpha_{e,1}$	$\alpha_{e,2}$	$r_{b;1}$	$r_{b;2}$	α	$U^{(1)}$	<i>U</i> ⁽²⁾
0.1	0.9	13.37	13.37	0.01	0.01	24.73	64.82	0.06	-0.48	-0.50
0.2	0.8	13.37	13.37	0.01	0.01	26.20	48.92	0.06	-1.23	-1.24
0.3	0.7	13.37	13.37	0.01	0.01	25.39	37.56	0.05	-0.94	-0.94
0.4	0.6	13.37	13.37	0.01	0.01	29.94	36.09	0.06	-0.90	-0.90
0.5	0.5	13.37	13.37	0.01	0.01	18.71	18.71	0.02	-0.98	-0.98
0.6	0.4	13.37	13.37	0.01	0.01	36.09	29.94	0.06	-0.90	-0.90
0.7	0.3	13.37	13.37	0.01	0.01	37.56	25.39	0.05	-0.94	-0.94
0.8	0.2	13.37	13.37	0.01	0.01	48.92	26.20	0.06	-1.24	-1.23
0.9	0.1	13.37	13.37	0.01	0.01	64.82	24.73	0.06	-0.50	-0.48
1	9	6.20	6.20	0.03	0.03	10.30	25.74	0.10	-2.02	-2.06
2	8	6.20	6.20	0.02	0.02	10.92	19.79	0.10	-1.50	-1.54
3	7	6.20	6.20	0.02	0.02	11.66	16.81	0.10	-1.28	-1.29
4	6	6.20	6.20	0.02	0.02	12.47	14.89	0.10	-1.13	-1.13
5	5	6.20	6.20	0.02	0.02	7.32	7.32	0.03	-1.14	-1.14
6	4	6.20	6.20	0.02	0.02	14.89	12.47	0.10	-1.13	-1.13
7	3	6.20	6.20	0.02	0.02	16.81	11.66	0.10	-1.29	-1.28
8	2	6.20	6.20	0.02	0.02	19.79	10.92	0.10	-1.54	-1.50
9	1	6.20	6.20	0.02	0.02	25.74	10.30	0.10	-2.06	-2.02
10	90	2.88	2.88	0.06	0.06	4.32	10.11	0.16	-2.28	-2.39
20	80	2.88	2.88	0.06	0.06	4.41	7.72	0.15	-1.63	-1.73
30	70	2.88	2.88	0.06	0.06	4.66	6.62	0.15	-1.43	-1.46
40	60	2.88	2.88	0.06	0.06	5.01	5.93	0.15	-1.28	-1.29
50	50	2.88	2.88	0.06	0.06	3.08	3.08	0.05	-1.34	-1.34
60	40	2.88	2.88	0.06	0.06	5.93	5.01	0.15	-1.29	-1.28
70	30	2.88	2.88	0.06	0.06	6.62	4.66	0.15	-1.46	-1.43
80	20	2.88	2.88	0.06	0.06	7.72	4.41	0.15	-1.73	-1.63
90	10	2.88	2.88	0.06	0.06	10.11	4.32	0.16	-2.39	-2.28
100	900	1.34	1.34	0.12	0.12	1.82	3.96	0.24	-2.02	-2.29
200	800	1.34	1.34	0.12	0.12	1.78	2.98	0.21	-1.43	-1.61
300	700	1.34	1.34	0.12	0.12	1.92	2.66	0.22	-1.34	-1.41
400	600	1.34	1.34	0.12	0.12	1.95	2.29	0.20	-2.97	-3.00
500	500	1.34	1.34	0.12	0.12	1.39	1.39	0.09	-1.36	-1.36
600	400	1.34	1.34	0.12	0.12	2.29	1.95	0.20	-3.00	-2.97
700	300	1.34	1.34	0.12	0.12	2.66	1.92	0.22	-1.41	-1.34
800	200	1.34	1.34	0.12	0.12	2.98	1.78	0.21	-1.61	-1.43
900	100	1.34	1.34	0.12	0.12	3.96	1.82	0.24	-2.29	-2.02

Table 2. The same as in Table 1 but for the case of $Z_1 = Z_2 = 1$.

6. Results and Discussions

6.1. The Properties of the Obtained Solutions

As a continuation of our previous research [1,2], in this work, fully ionized electron–ion plasmas are chosen with the positive ion charges of two different kinds. Such a choice is especially important, as increasing the number of ion components further would not cause the appearance of any new phenomena.

One can see that the procedures of obtaining Equations (17)–(33), as well as the values of the existing parameters, provide that these expressions are self-consistent; satisfy all the conditions from Section 2.1, including Equations (37) and (39); and can be applied not only to the classical but also to the quantum-mechanical systems (see [2]), including here the plasmas of higher non-ideality. Because the presented expressions do not contain the particle masses, they can also be used for describing some other systems (the corresponding electrolytes and dusty plasmas). The behaviour of the ion and electron densities is illustrated in Figures 2 and 3 for the examples of the cases of (*i*1) and (*i*2) for $Z_2 \neq Z_1$ and $Z_2 = Z_1$, respectively.



Figure 2. The behavior of reduced densities $n_1^{(1)}(r)/N_1$ (curve marked with 1), $n_2^{(1)}(r)/N_2$ (curve marked with 2), $n_2^{(2)}(r)/N_2$ (curve marked with 3), $n_1^{(2)}(r)/N_1$ (curve marked with 4), and $n_{1,2}^{(e)}(r)/N_e$ (curve marked with 5 and 6) in the case of $Z_1 = 1$, $Z_2 = 1$ and $T_i = T_e = T$, where T = 30,000 K.



Figure 3. The behavior of reduced densities $n_1^{(1)}(r)/N_1$ (curve marked with 1), $n_2^{(1)}(r)/N_2$ (curve marked with 2), $n_2^{(2)}(r)/N_2$ (curve marked with 3), $n_1^{(2)}(r)/N_1$ (curve marked with 4), and $n_{1,2}^{(e)}(r)/N_e$ (curve marked with 5 and 6), in the case of $Z_1 = 1$, $Z_2 = 2$ and $T_i = T_e = T$, where T = 30,000 K.

Because Equations (27) and (32) show that the solutions $n_e^{(1,2)}(r)$ and $n_{i;1,2}^{(i)}(r)$ are singular at the point r = 0, it is useful to note that the existence of singularities in model solutions is fully acceptable, if it does not have other non-physical consequences. Such solutions are well known in physics; it is enough to mention, for example, the Thomas–Fermi models of electron shells of heavy atoms ([37,38]; see also [39]), which have been used in plasma research up to the present (see e.g., [40]). Except for the potential $\Phi^{(1,2,e)}(r)$ and $\varphi^{(1,2,e)}$, the systems $S_a^{(1,2,e)}$ are certainly characterized by radial charge densities $P^{(1,2,e)}(r) \equiv 4\pi r^2 \times \rho^{(1,2,e)}(r)$. According to [3], each of the functions $|P^{(1,2,e)}(r)|$ has at least one strongly expressed maximum, whose position is an important characteristic of the distribution of charge in the neighborhood of the probe particle.

In order to demonstrate the very large differences between the alternative and DH-like characteristics, we compare the asymptotic behaviour of the potential $\Phi^{(1,2,e)}(r)$ and DH-like potential $\Phi^{(1,2,e)}_{DH}(r)$:

$$\Phi^{(1,2)}(r) \sim Z_{1,2}e \times \frac{e^{-\kappa_{as;1,2}(r-r_{b;1,2})}}{r}, \quad r > r_{b;1,2}, \quad \Phi^{(1,2,e)}_{DH}(r) \sim Z_{1,2,e}e \times \frac{e^{-\kappa_{DH}r}}{r}, \quad r > 0$$
(42)

where the ion screening constants $\kappa_{as;1,2}$ are given by Equation (24) and the DH screening constant $\kappa_{DH} = (\kappa_{0;1}^2 + \kappa_{0;2}^2 + \kappa_{0;e}^2)^{1/2}$, where $\kappa_{0;1,2}$ and $\kappa_{0;e}$ are determined by Equations (24) and (29). Here, we consider the case of classical plasma with $T_i = T_e = T$, where $\partial \mu_{1,2,e} / \partial N_{1,2,e} = kT / N_{1,2,e}$ and, consequently, the relations

$$\frac{\kappa_{as;1,2}}{\kappa_{DH}} \equiv \frac{r_{DH}}{r_{as;1,2}} = \frac{[Z_{1,2}^2 N_{1,2} (1 - \alpha_{e;1,2}) (1 - \alpha_i)]^{\frac{1}{2}}}{(Z_1^2 N_1 + Z_2^2 N_2 + N_e)^{\frac{1}{2}}}$$
(43)

are valid. These relations show that the ion asymptotic screening constants $\kappa_{as;1,2}$ always have to be significantly smaller than κ_{DH} , and at the same time, the corresponding screening radii $r_{as;1,2}$ always have to be significantly larger then r_{DH} . It is important that a similar result obtained in [2] was noted there as an evident shortcoming of the DH solution.

From Equation (43), it follows that $\Phi^{(1,2,e)}(r)$ has a completely different asymptotic behavior compared to $\Phi_{DH}^{(1,2,e)}(r)$. We note that we reach the same conclusion by comparing the behavior of the radial charge density $\rho^{(1,2,e)}(r)$ to its DH-like analog.

As the main characteristics of the considered plasmas we take here, the probe particle mean potential energies $U^{(1,2,e)}$ are later identified with the mean potential energies of ions in the real plasmas. In order to determine these ion energies $U^{(1,2,e)}$, it is necessary to know the values of the potential $\varphi^{(1,2,e)}$:

$$\varphi^{(1,2,e)} = \int_0^\infty \frac{\rho^{(1,2,e)}(r)}{r} 4\pi r^2 dr = 4\pi \int_0^\infty \rho^{(1,2,e)}(r) r dr \tag{44}$$

where $\rho^{(1,2,e)}(r)$ denotes the charge density. We recall that the case $Z_1 = Z_2 = 1$ can correspond to the case of plasma with the ion H⁺ or He⁺(1s), and so forth. Within this work, the energies $U^{(1)}$ and $U^{(2)}$ are determined for two cases: $Z_1 = 1$ and $Z_2 = 2$, and $Z_1 = Z_2 = 1$. In both cases, the calculations are performed for plasmas with the electron densities $10^{16} \text{ cm}^{-3} \le N_e \le 10^{20} \text{ cm}^{-3}$, for $T = 3 \times 10^4 \text{ K}$. The obtained results are presented in Tables 1 and 2 with fairly small ion-density steps. In order to investigate the dependance of the energies of the systems on the temperature, the calculations of $U^{(1)}$ and $U^{(2)}$ were performed and are presented in Tables 3 and 4 for $N_e = 10^{19} \text{ cm}^{-3}$ and for the temperatures $T = 1 \times 10^4$, 1.5×10^4 , 2×10^4 and 2.5×10^4 K. From these results, one can see that the potential energies $U^{(1)}$ and $U^{(2)}$ are sensitive to a considerable lowering of the temperature ($T = 1 \times 10^4$ K).

Table 3. The potential energies $U^{(1)}$ (in eV) and $U^{(2)}$ (in eV) for the case of $Z_1 = 1$ and $Z_2 = 2$ at $N_e = 10^{19}$ cm⁻³ and $T = 1 \times 10^4$, 1.5×10^4 , 2.0×10^4 , and 2.5×10^4 K. The densities $N_{1,2}$ are in 10^{18} cm⁻³.

N_1	N_2	<i>U</i> ⁽¹⁾	<i>U</i> ⁽²⁾						
		10,0	00 K	15,0	00 K	20,0	00 K	25,0	00 K
0.5	4.75	-1.98	-1.53	-1.00	-3.90	-1.53	-1.71	-1.93	-2.16
1.0	4.50	-1.69	-1.33	-0.97	-3.51	-3.94	-3.86	-4.70	-4.71
1.5	4.25	-1.37	-1.15	-3.17	-2.93	-3.32	-3.36	-3.88	-4.01
2.0	4.00	-1.33	-1.11	-1.05	-3.59	-1.97	-1.95	-3.42	-3.62
2.5	3.75	-1.15	-1.01	-1.11	-3.97	-2.49	-2.72	-3.13	-3.37
3.0	3.50	-1.20	-1.04	-1.13	-3.99	-2.44	-2.61	-3.02	-3.22
3.5	3.25	-1.35	-1.12	-1.91	-1.96	-2.34	-2.50	-2.94	-3.11
4.0	3.00	-1.19	-1.04	-1.16	-4.16	-2.37	-2.50	-2.91	-3.06
4.5	2.75	-1.14	-1.29	-2.34	-2.24	-2.36	-2.47	-2.78	-2.92
5.0	2.50	-1.37	-1.17	-2.52	-2.38	-2.57	-2.62	-2.83	-2.94
5.5	2.25	-1.05	-1.21	-2.44	-2.32	-4.02	-3.83	-2.77	-2.87
6.0	2.00	-1.08	-1.27	-2.32	-2.24	-4.32	-4.09	-3.78	-3.74
6.5	1.75	-1.64	-1.39	-2.39	-2.30	-3.48	-3.38	-1.89	-6.48
7.0	1.50	-1.01	-0.99	-1.86	-1.51	-2.43	-2.18	-2.92	-2.84
7.5	1.25	-1.20	-1.28	-2.09	-1.71	-2.73	-2.44	-3.62	-3.51
8.0	1.00	-1.54	-1.85	-2.55	-2.12	-3.61	-3.42	-2.14	-4.98
8.5	0.75	-2.03	-1.32	-1.32	-3.01	-1.93	-2.10	-2.56	-2.89
9.0	0.50	-0.83	-1.94	-1.62	-1.59	-2.41	-2.55	-3.11	-2.70
9.5	0.25	-1.20	-1.00	-2.36	-2.74	-3.37	-2.83	-4.38	-3.85

Table 4. The same as in Table 3 but for the case of $Z_1 = Z_2 = 1$.

N_1	N_2	$U^{(1)}$	<i>U</i> ⁽²⁾						
		10,0	00 K	15,0	00 K	20,0	00 K	25,0	00 K
0.5	9.50	-0.82	-0.94	-1.50	-1.62	-2.13	-2.26	-0.76	-0.87
1.0	9.00	-0.52	-0.65	-1.04	-1.16	-1.49	-1.61	-1.91	-2.03
1.5	8.50	-0.39	-0.52	-0.84	-0.96	-1.21	-1.33	-1.56	-1.67
2.0	8.00	-0.39	-0.47	-0.76	-0.84	-1.07	-1.15	-1.38	-1.49
2.5	7.50	-0.37	-0.42	-0.74	-0.79	-1.00	-1.05	-1.30	-1.35
3.0	7.00	-0.35	-0.38	-0.66	-0.70	-0.97	-1.01	-1.18	-1.22
3.5	6.50	-1.03	-1.05	-0.65	-0.67	-0.92	-0.95	-1.16	-1.18
4.0	6.00	-0.94	-0.95	-0.64	-0.65	-0.85	-0.87	-1.06	-1.08
4.5	5.50	-0.95	-0.95	-1.50	-1.51	-0.82	-0.83	-1.05	-1.06
5.0	5.00	-0.34	-0.34	-0.66	-0.66	-0.95	-0.95	-1.18	-1.18
5.5	4.50	-0.95	-0.95	-1.51	-1.50	-0.83	-0.82	-1.06	-1.05
6.0	4.00	-0.95	-0.94	-0.65	-0.64	-0.87	-0.85	-1.08	-1.06
6.5	3.50	-1.05	-1.03	-0.67	-0.65	-0.95	-0.92	-1.18	-1.16
7.0	3.00	-0.38	-0.35	-0.70	-0.66	-1.01	-0.97	-1.22	-1.18
7.5	2.50	-0.42	-0.37	-0.79	-0.74	-1.05	-1.00	-1.35	-1.30
8.0	2.00	-0.47	-0.39	-0.84	-0.76	-1.15	-1.07	-1.49	-1.38
8.5	1.50	-0.52	-0.39	-0.96	-0.84	-1.33	-1.21	-1.67	-1.56
9.0	1.00	-0.65	-0.52	-1.16	-1.04	-1.61	-1.49	-2.03	-1.91
9.5	0.50	-0.94	-0.82	-1.62	-1.50	-2.26	-2.13	-0.87	-0.76

6.2. Interpretation of the Obtained Results

Results such as those presented in the previous section (see Equation (43)) might leave an impression of having an absolute advantage over the DH-like methods (or other similar methods). Such an impression is incorrect, as an absolute advantage of the presented method is for the case of a system of the closed type. In the same context, it is necessary to interpret the phenomena that were described in the introduction concerning the results presented in the Figure 1a,b; these phenomena can be interpreted as physically unacceptable when the DH or DH-like methods are used on the system of the closed type. However, the system receives treatment as closed-type only in the case in which it is described by means of the above-mentioned additional conditions. Consequently, in the opposite case (when additional conditions are absent), the system can be successfully described by means of DH-like or similar methods. Concerning this, we refer to Figure 4a,b, which shows the results of the application of the DH method to the considered plasma ((a) $Z_1 = 1$ and $Z_2 = 2$, and (b) $Z_1 = Z_2 = 1$). It is useful to compare this figure with Figures 2 and 3. In this context, we draw attention to the fact that such a treatment itself has to be determined on the basis of the properties of the considered system and the physical problem, which can be solved using that system.

At the end of this point, it would be useful to linger on such influence of the additional conditions on the properties of the obtained solutions, which could be treated as a manifestation of their deviation from thermodynamic equilibrium. Here, we refer to the necessity of substituting the equation obtained from the condition of thermodynamic equilibrium by the equations obtained in different ways—the electron charge density $n_e^{(1,2)}$ in the region of large r, and the ion charge density $n_{1,2}^{(2,1)}(r)$, also in the region of large r. However, we draw attention to the fact that all changes have purely phenomenological characteristics and do not influence the thermodynamic properties of the considered gases: gas with temperature T remains a gas with the temperature T. Additionally, we recall the fact that in all regions of r, where the considered components can be treated independently from one another, their state was described by means of equations obtained from the condition of thermodynamic equilibrium.

From the above-presented material, it follows that the basic model can generate only the DH or some DH-like methods. In this sense, this model has already exhausted its potential, but as one can see, it enables, with minimal deviation from the basic model (in the area of mathematical apparatus), us to leave the DH-like sphere and develop a new model method of describing the plasma's inner electrostatic screening.

6.3. The Possible Ion–Ion Probe Systems

From the presented work, it follows that the main properties of the considered three-component system originate from the analogy with the properties of a positron–ion probe system. Concerning this, it is useful to note the fact that was established by means the molecular dynamic (MD) simulation of a dense electron-proton plasma in [41]. In this work, some characteristics of the considered plasma were determined as the results of averaging over all ion configurations possible under the considered conditions, for different values of the ratio m_e/m_p , where m_e and m_p are the electron and proton masses. These values were changed from 1/1836 to 1/100, but the changes of the results of MD simulations could be neglected. This can be very interesting, even only for the similarity of the procedures that were used here and in [41]. However, it can be particularly important under the assumption that a similar conclusion is valid in the case of plasma that contains electrons and ions of some heavy atoms, particularly if the non-negligible probability is taken into account, meaning that the value of the mentioned ratio can be even greater than 1/100.



Figure 4. (a) The reduced Debye-Hückel (DH) densities $n_{D;1}^{(1)}(r)/N_1$ (curve marked with 1), $n_{D;2}^{(1)}(r)/N_2$ (curve marked with 2) and $n_{D;e}^{(1)}(r)/N_e$ (curve marked with 3) in the case of $Z_1 = 1$, $Z_2 = 2$ and $T_i = T_e = T$, where T = 30,000 K. (b) The reduced DH densities $n_{D;1}^{(1)}(r)/N_1$ (curve marked with 1), $n_{D;2}^{(1)}(r)/N_2$ (curve marked with 2), and $n_{D;e}^{(1)}(r)/N_e$ (curve marked with 3) in the case of $Z_1 = Z_2 = 1$ and $T_i = T_e = T$, where T = 30,000 K.

7. Conclusions

The object of the investigation of fully ionized electron-ion plasma was chosen with positively charged ions of two different kinds, including here the plasmas of higher non-ideality. Within the presented method, such extremely significant phenomena as the electron-ion and ion-ion correlations are included. The collective effect of many correlated particle interactions strongly affects the spectral properties of atoms and properties of their collision processes with respect to those for isolated systems. The screening characteristics of the considered plasmas in a wide region of the electron densities and temperatures have been calculated. Here, the case of the three-component system was considered, which is especially important as we expect that a further increase in the number of ion components in the more complicated systems would not cause the appearance of any new phenomena.

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The Application of the Cut-Off Coulomb Model Potential for the Calculation of Bound-Bound State Transitions

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Abstract: In this contribution, we present results of bound state transition modeling using the cut-off Coulomb model potential. The cut-off Coulomb potential has proven itself as a model potential for the dense hydrogen plasma. The main aim of our investigation include further steps of improvement of the usage of model potential. The results deal with partially ionized dense hydrogen plasma. The presented results covers $N_e = 6.5 \times 10^{18} \text{ cm}^{-3}$, T = 18,000 K and $N_e = 1.5 \times 10^{19} \text{ cm}^{-3}$, T = 23,000 K, where the comparison with the experimental data should take place, and the theoretical values for comparison. Since the model was successfully applied on continuous photoabsorption of dense hydrogen plasma in the broad area of temperatures and densities, it is expected to combine both continuous and bound-bound photoabsorption within single quantum mechanical model with the same success.

Keywords: atomic processes; radiative transfer; Sun: atmosphere; Sun: photosphere; stars: atmospheres; white dwarfs

1. Introduction

The problems of plasma opacity, energy transport and radiative transfer under moderate and strong non-ideality are of interest in theoretical and experimental research [1–4]. The strong coupling and density effects in plasma radiation were the subject of numerous experimental and theoretical studies in the last decades. Here, we keep in mind the plasma of the inner layers of the solar atmosphere, as well as of partially ionized layers of other stellar atmospheres—for example, the atmospheres of DA white dwarfs with effective temperatures between 4500 K and 30,000 K.

In this paper, we presented a new model way of describing atomic photo-absorption processes in dense, strongly ionized hydrogen plasmas, which is based on the approximation of the cut-off Coulomb potential. By now, this approximation has been used in order to describe transport properties of dense plasmas (e.g., [1,5,6]), but it was clear that it could be applied to some absorption processes in non-ideal plasmas too [3,7–9]. This topic itself, and the search for more consistent models of screening and more realistic potentials in plasmas is still continuing and is very real (e.g., [10–13] and references therein).

The neutral hydrogen acts as an absorber in plasma, and its concentration determination is essential in order to calculate the absorption coefficients. The neutral hydrogen concentration from experimental data from [14] is used, and, since the result presented here is in the area where the Saha equation is valid, the neutral concentrations using the Saha equation with the ionization potential correction is possible. Since this is related to the further step, bound level broadening mechanism,

the procedure was not elaborated here. In addition, the emitter, neutral hydrogen, interacts with other plasma species such as electrons and ions. In such a way, the theoretical hydrogen plasma model results as well as NIST database values act as a limiting case for verification of solutions, in the case when plasma influence is small or diminishes, e.g., $r_c \rightarrow \infty$ in later case.

In order to calculate the total absorption coefficient, the same broadening mechanism for the bound state levels should be applied. Up until now, the continuous absorption processes were calculated with the help of parametric broadening for the bound-free absorption [7]. The broadening processes play a more important role in bound-bound processes, and, as such, the more realistic theoretical model of broadening of bound state levels or a result of adequate molecular dynamics (MD) simulations should be used. Only in such a case could the total absorption coefficients, obtained within the same broadening model, be compared with real experimental data. This work is in progress.

A first step in extending of the model with additional processes is the bound state transition processes inclusion. The bound state transition processes are stated as the most important goal in the development of the self containing model, capable of describing optical as well as dynamical characteristics of dense hydrogen plasma. The characteristics of the bound state transitions in plasma diagnostics are well known, an almost mandatory method [15]. The usage of the hydrogen lines as a probing method for the plasma characteristics is also well known and widely used in plasmas of moderate and small non-ideality [16]. In accordance with that previously mentioned, the continuation of research of the presented approach is the inclusion of the bound-bound photo absorption processes:

$$\varepsilon_{\lambda} + \mathrm{H}^{*}(n_{i}, l_{i}) \to \mathrm{H}^{*}(n_{f}, l_{f}), \tag{1}$$

where *n* and *l* are the principal and the orbital quantum number of hydrogen-atom excited states, hydrogen atom in its initial state $|n_i, l_i >$ is presented by $H^*(n_i, l_i)$, its final state $|n_f, l_f >$ by $H^*(n_f, l_f)$, and ε_{λ} presents absorbed photon energy.

2. Theory

2.1. The Approximation of the Cut-Off Coulomb Potential

The absorption processes (1) in astrophysical plasma are considered here as a result of radiative transition in the whole system "electron–ion pair (atom) + the neighborhood", namely: $\varepsilon_{\lambda} + (H^+ + e)_i + S_{rest} \rightarrow (H^+ + e)_f + S'_{rest}$, where S_{rest} and S'_{rest} denote the rest of the considered plasma. However, as it is well known, many-body processes can sometimes be simplified by their transformation to the corresponding single-particle processes in an adequately chosen model potential.

As an adequate model potential for hydrogen plasma with such density, we choose, as in [3,5], the screening cut-off Coulomb potential, which satisfies above conditions, and can be presented in the form:

$$U(r;r_c) = \begin{cases} -\frac{e^2}{r}, & 0 < r \le r_c, \\ U_{p;c}, & r_c < r < \infty, \end{cases}$$
(2)

where the cut-off radius r_c is defined by relation (3), as it is illustrated by Figure 1a. Here, e is the modulus of the electron charge, r—distance from the ion, and cut-off radius r_c —the characteristic screening length of the considered plasma. Namely, within this model, it is assumed that quantity

$$U_{p;c} = -\frac{e^2}{r_c} \tag{3}$$

is the mean potential energy of an electron in the considered hydrogen plasma.



Figure 1. Behaviour of the potentials $U(r; r_c)$ and $U_c(r)$, where r_c is the cut-off parameter. In (**a**) the potential goes to the mean potential energy of an electron in the considered hydrogen plasma $U_{p;c}$, and in (**b**), the value of $U_{p;c}$ is taken as zero energy, i.e., $U_c = U(r; r_c) - U_{p;c}$, in the more applicable form used here.

As in [3,5,8], we will take the value $U_{p;c}$ as the zero of energy. After that, the potential Equation (2) is transformed to the form

$$U_{c}(r) = \begin{cases} -\frac{e^{2}}{r} + \frac{e^{2}}{r_{c}}, & 0 < r \le r_{c}, \\ 0, & r_{c} < r < \infty, \end{cases}$$
(4)

which is illustrated by Figure 1b. Here, e is the modulus of the electron charge, r—distance from the ion, and cut-off radius r_c —the characteristic screening length of the considered plasma.

It is important that the cut-off radius r_c can be determined as a given function of N_e and T, using two characteristic lengths:

$$r_i = \left(\frac{k_B T}{4\pi N_i e^2}\right)^{1/2} \quad r_{s;i} = \left(\frac{3}{4\pi N_i}\right)^{\frac{1}{3}},\tag{5}$$

namely, taking that $N_i = N_e$ and $r_c = a_{c;i} \cdot r_i$ we can directly determine the factor $a_{c;i}$ as a function of ratio $r_{s;i}/r_i$, on the basis of the data about the mean potential energy of the electron in the single ionized plasma from [12]. The behavior of $a_{c;i}$ in a wide region of values of $r_{s;i}/r_i$ is presented in Figure 2.



Figure 2. Behavior of the parameters $a_{c;i} \equiv r_c/r_i$ as the function of the ratio $r_{s;i}/r_i$, where r_i is given by Equation (5) and $r_{s;i}$ is the ion Wigner–Seitz radius for the considered electron-ion plasma (5). The presented curve is obtained on the basis of data presented in [12].

2.2. The Calculated Quantities

In accordance with that, the behavior of the dipole matrix element is investigated here. It is given by

$$\hat{D}(r; r_c; n_i, l_i; n_f, l_f) = < n_f, l_f |\mathbf{r}| n_i, l_i >,$$
(6)

where the wave functions $|n_i, l_i\rangle$ and $|n_f, l_f\rangle$ are initial and final state wave functions obtained within the model of cut-off Coulomb potential, for the calculations of the plasma characteristics, or the theoretical hydrogen ones in order to additionally check the model.

For the calculation of oscillator strength, we use expressions from [17–19]

$$f(n_f, l_f; n_i, l_i; r_c) = \frac{1}{3} \frac{\nu}{Ry} \left[\frac{max(l_f, l_i)}{2l_f + 1} \right] \hat{D}(r; r_c; n_i, l_i; n_f, l_f)^2,$$
(7)

where *Ry* is the Rydberg constant, in the same units as the frequency ν of the transition $(n, l) \rightarrow (n', l')$. The calculated data are presented in Table 1.

Our future plan is to include broadening processes and make it possible to directly compare data with the experimental ones (e.g., the data from [14]). In continuation, some of the bound-bound theoretical aspects are shown.

The total absorption cross section of the line could be linked with the dipole moment directly with the help of relation

$$\sigma_0(\omega = \omega_{fi}) = \frac{1}{3} \frac{g_2}{g_1} \frac{\pi \omega_{fi}}{\epsilon_0 \hbar c} \hat{D}(r; r_c; n_i, l_i; n_f, l_f)^2.$$
(8)

It should be noted that the absorption line profile is needed in order to apply the obtained results to the absorption calculations:

$$\sigma(\omega) = \sigma_0(\omega = \omega_{fi})g(\omega, \omega_{fi}).$$
(9)

Here, $g(\omega, \omega_{fi})$ is the spectral line profile and could be obtained, for instance, as a result of a molecular dynamics (MD) simulations [20–22] or as a result of theoretical modeling and spectral line data [23–27]. The lineshape is normalized to unity area, e.g.,

$$\int_{-\infty}^{\infty} g(\omega) d\omega = 1.$$
 (10)

The line profile is a convolution of the broadening of the initial and final state energy levels, e.g.,

$$g(\omega) = g_i(\omega) * g_f(\omega). \tag{11}$$

Here, g_i and g_f are energy level broadening shapes of the initial and final states. From here, the absorption coefficient could be calculated

$$k(\omega) = N_i \int_f \int_i g_i(\omega) g_f(\omega) \sigma_0.$$
(12)

Here, N_i is the concentration of hydrogen in initial level $|n_i, l_i >$. The broadening of the levels should be considered separately in order to use the same broadening profiles for the calculation of the bound-free transition continuous photo absorption.

There are a few steps to be carried out, and the first to be taken into account is to use a parametric level and line broadening model. The second one is to use a theoretical model for spectral line broadening and try to deconvolve level broadening values, and the final step should be the usage of the MD code [20–22]. A candidate for the coupling of the MD code is LAMMPS, which is capable of simulating a large scale molecular dynamics simulations, and it also could include particular interaction potential as well as all relevant processes of interest not included in the initial code as

an additional module. The broadening parameters could be calculated from average energy and particle distributions and their temporal distributions, or, as an iterative procedure, a more real plasma potential could be obtained [20,21,28,29]. In the former case, the solutions would not necessary to be described as a set of analytical functions. Since there are more steps, we did not elaborate on the ideas until we obtained some initial results.

Table 1. The oscillator strength values for the "short", $Ne = 1.5 \times 10^{19}$ cm⁻³, T = 23,000 K, and "long" pulse, $Ne = 6.5 \times 10^{18}$ cm⁻³, T = 18,000 K, from [14], as well as a theoretical hydrogen case calculated by code from [18], and NIST spectral database values [30].

		Short Pulse	Long Pulse	Theory	NIST
$ n_i, l_i >$	$ n_f, l_f >$	$f(n_f, l_f; n_i, l_i; r_c)$	$f(n_f, l_f; n_i, l_i; r_c)$	$f(n_f, l_f; n_i, l_i)$	$f(n_f, l_f; n_i, l_i)$
1,0>	2,1>	0.416197	0.416197	0.416200	0.416400
1,0>	3,1>	0.079102	0.079102	0.079101	0.079120
1,0>	4,1>	0.028923	0.028991	0.028991	0.029010
1,0>	5,1>		0.013728	0.013938	0.013950
2,0>	3,1>	0.434865	0.434865	0.434870	0.435100
2,0>	4,1>	0.102533	0.102756	0.102760	0.102800
2,0>	5,1>		0.041425	0.041930	0.041950
2,1>	3,0>	0.013589	0.013589	0.013590	0.013600
2,1>	3,2>	0.695785	0.695785	0.695780	0.696100
2,1>	4,0>	0.003035	0.003045	0.003045	0.003046
2,1>	4,2>	0.121659	0.121795	0.121800	0.102800
2,1>	5,0>		0.001191	0.001213	0.001214
2,1>	5,2>		0.043962	0.044371	0.044400
3,0>	4,1>	0.483750	0.484708	0.484710	0.484900
3,0>	5,1>		0.119310	0.121020	0.121100
3,1>	4,0>	0.032165	0.032250	0.032250	0.032280
3,1>	4,2>	0.617675	0.618282	0.618290	0.618600
3,1>	5,0>		0.007299	0.007428	0.007433
3,1>	5,2>		0.138013	0.139230	0.139300
3,2>	4,1>	0.010971	0.010992	0.010992	0.011000
3,2>	4,3>	1.017260	1.017520	1.017500	1.018000
3,2>	5,1>		0.002180	0.002210	0.002211
3,2>	5,3>		0.156046	0.156640	0.156700
4,0>	5,1>		0.537527	0.544150	0.544400
4,1>	5,0>		0.052123	0.052907	0.052940
4,1>	5,2>		0.604678	0.609290	0.609700
4,2>	5,1>		0.027498	0.027822	0.027840
4,2>	5,3>		0.887328	0.890250	0.890600
4,3>	5,2>		0.008809	0.008871	0.008877
4,3>	5,4>		1.344790	1.345800	1.346000

3. Results and Discussion

The convergence toward theoretical values for hydrogen is examined and proven for the bound-bound transitions that could appear in the investigated area of electron concentrations and temperatures, [31]. Since the initial results have proven that the model potential could be used for the bound-bound state transition calculations, further investigation was completed.

The next step towards the application of the model results is a calculation of the oscillator strength values for the plasma parameters denoted as "short", with $N_e = 1.5 \times 10^{19}$ cm⁻³, T = 23,000 K and "long" pulse, with $N_e = 6.5 \times 10^{18}$ cm⁻³, T = 18,000 K from [14]. The radius r_i , given by Equation (5), Wigner–Seitz and cut-off radius as well as atom concentrations given by the Saha equation are $r_i = 51.0655$ a.u., $r_{s;i} = 47.5339$ a.u., $r_c = 44.9907$ a.u., $N_a = 1.9 \times 10^{19}$ cm⁻³ for short and $r_i = 68.6260$ a.u., $r_{s;i} = 62.8149$ a.u., $r_c = 60.4062$ a.u., $N_a = 3.4 \times 10^{19}$ cm⁻³ for long pulse.

The essential quality of the presented approach is that the values are calculated with the help of wave functions that are obtained as a completely analytical solution in the frame of the presented model. The influence of the numerical procedure is minimized, and, as a consequence, the possibility of introducing the numerical artifacts into the solution is also minimized.

Since the convergence towards the theoretical, unperturbed Coulomb potential is investigated in [31], and the results of the presented calculations converge uniformly towards theoretical values, it is expected that the presented model could obtain good data in a wider variety of plasma parameters than shown here.

There is a need to introduce the broadening mechanism for the energy levels in the presented approach. There is a possibility to couple the presented model calculations with the MD code in order to obtain a consistent model for the broadening of the bound levels. Those results influence the bound-free and bound-bound absorption profiles and are essential for the hydrogen plasma optical properties calculation in the frame of the cut-off Coulomb potential model approach.

4. Conclusions

The presented results are a step forward towards inclusion of the entire photo-absorption processes for hydrogen atoms in plasma within the frame of the cut-off Coulomb potential model. One of the benefits of the presented results is a completely quantum mechanical solution for the cut-off Coulomb model potential, obtained from wave functions that are analytical and represented with the help of special functions, e.g., the influence of additional numerical source of errors is minimized as much as possible. Along with this, the solutions converge towards theoretical pure Coulomb potential ones as the cut-off radius converges to infinity, e.g., the influence of plasma diminishes. Further steps in application of the presented results, the inclusion of the bound-bound transition processes within the continuous absorption model, as well as its application in the analysis of the spectral absorption processes is to be carried out. In order to make such calculations, it is needed to include the models for the bound state level broadening. Our plan is to present the results obtained during this investigation in a database that can be accessed directly through http://servo.aob.rs as a web service, similar to the existing databases [26,27,32].

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Author Contributions: N. M. S. have adopted the cut-off Coulomb potential model for the optical plasma characteistics; N. M. S. developed the code and performed the calculations; M. S. D., Z. J. S. and V. A. S. analysed the data; N. M. S., V. A. S. and M. S. D. wrote the paper; M. S. D., V. A. S. and Z. J. S. have edited the paper.

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Article Detailed Investigation of the Electric Discharge Plasma between Copper Electrodes Immersed into Water

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Abstract: A phenomenological picture of pulsed electrical discharge in water is produced by combining electrical, spectroscopic, and imaging methods. The discharge is generated by applying ~350 μ s long 100 to 220 V pulses (values of current from 400 to 1000 A, respectively) between the point-to-point copper electrodes submerged into the non-purified tap water. Plasma channel and gas bubble occur between the tips of the electrodes, which are initially in contact with each other. The study includes detailed experimental investigation of plasma parameters of such discharge using the correlation between time-resolved high-speed imaging, electrical characteristics, and optical emission spectroscopic data. Radial distributions of the electron density of plasma is estimated from the analysis of profiles and widths of registered H_{α} and H_{β} hydrogen lines, and Cu I 515.3 nm line, exposed to the Stark mechanism of spectral lines' broadening. Estimations of the electrodes' erosion rate and bubbles' size depending on the electrical input parameters of the circuit are presented. Experimental results of this work may be valuable for the advancement of modeling and the theoretical understanding of the pulse electric discharges in water.

Keywords: laboratory plasma; spectra; line profiles

1. Introduction

There is increasing interest in plasma discharge in liquid, mostly because of its importance in electrical transmission processes and its practical applications in biology, chemistry, and electrochemistry. Special place within the variety of its exploitations belongs to the water treatment. Due to the low efficiency of the conventional techniques, and presence of a number of disadvantages in other developing and existing methods (i.e., chlorination, ozonation, advanced oxidation processes, photocatalysis) [1–3], application of the electrical discharges in liquid has proven to be one of the most advanced and affordable methods not only for the water treatment (removal of organic compounds), but also in surface treatment and plasma sterilizations (inactivation or killing of microorganisms) [4–6]. Such discharges are the effective sources of simultaneous production of intense UV radiation, shock waves, and various chemical products, including OH, O, HO₂, and H₂O₂ from the electric breakdown in water [7,8]. Moreover, shock waves produced by high-energy plasma discharges inside liquids are used for various applications, including underwater explosions [9], rock fragmentation [10], and lithotripsy [8]. A great deal of work has been done by Locke et al. [11], who presented the review of the current status of research on the application of high-voltage electrical discharges for promoting chemical reactions in the aqueous phase, with particular emphasis on applications to water cleaning. Another important application of the underwater electric discharges, which has attracted significant attention, is the nanomaterial synthesis by plasma-liquid interactions, including plasma-over-liquid and plasma-in-liquid configurations [12].

The nature of the discharges in liquids is much less understood and may be completely different from those for discharges in gases, therefore, in all the mentioned applications, it is important to understand the mechanism and dynamics of the electric breakdown process in liquids. Unfortunately, until now there are no complete physical models of underwater discharges, which makes it of a great scientific interest to investigate plasma discharges in liquid media.

This work, particularly, presents the phenomenological extension aiming to contribute to the better understanding of studies carried out in [13,14]. Investigations of nanoparticles interaction with biological environments are of great interest. It was found that colloidal substance is the most effective biological form of nanoparticles [15]. Moreover, it is known that solutions of silver and copper have bactericidal, antiviral, pronounced antifungal and antiseptic effects [16], therefore they are considered as perspective new biocides products. This partially explains the choice of the copper electrodes for the present study. Additionally, authors have worked on selection of copper spectral lines and corresponding spectroscopic data for diagnostics of multicomponent air plasma with copper admixtures in the past [17], therefore it is a reasonable material to start with.

The main motivation is to present a model describing physical processes occurring during electrospark dispersion of metal granules used for production of colloidal solutions, including the energy input calculations and studies of energy dissipation paths.

2. Experimental Method

2.1. Experimental Setup

The experimental setup is shown in Figure 1. It is composed of a pulsed generator (I), a trigger unit (II), a support (III) allowing positioning of the copper electrodes of 6 mm diameter (one is fixed and the other is movable) and a Pyrex container (IV) filled with non-purified water (pH ~6.5; t = 20-25 °C; conductivity ~5–50 mS/m). The current is measured with a Rogowsky coil (V), the data being stored with an oscilloscope (VI). The maximal load voltage U₀ is 430 V for a 1000 µF capacity and a maximal current of 4 kA.



Figure 1. Experimental setup for the underwater arc discharge generation.

Arc discharge is generated between the tips of two electrodes by applying a pulsed voltage of ~120, 150, 180 and 220 V, corresponding to the values of current 450, 660, 800 and 1000 A, respectively, with the average duration of the discharge of 320–360 μ s. The arc is triggered through the closing of a thyristor, with a switching time of about 100 μ s. The anode moves away when the arc ignites (due to repulsive electromagnetic forces).

The spectroscopic diagnostic is performed with an Acton SpectraPro SP-2750 (VII) spectrometer fitted with ProEM 1024 EMCCD camera (Princeton Instruments). The optical setup (Figure 1) is composed of a mirror (M1) and two lenses (L1 and L2), allowing to observe the arc from above. The arc is also visualized using a Photron Fastcam AX100 high-speed camera (VIII), the acquisition being synchronized to the arc triggering.

Discharge axis is perpendicular to the entrance slit of the spectrometer (opening width of 40 μ m). The apparatus function of the spectrometer is obtained with a low-pressure Hg lamp. Registration of the spectra is performed using 300 lines/mm grating (for H_{α} and Cu I lines with central wavelength set on $\lambda = 616.2$ and 515.0 nm, correspondingly) and 1800 lines/mm grating (for H_{β} line with central wavelength set on $\lambda = 486.1$ nm). The spectral resolution of the system is 0.056 nm for the 300 lines/mm grating and 0.0078 nm for 1800 lines/mm grating. The exposure time of the EMCCD camera is 200 μ s. It is triggered as the value of current rises above 120 A, which occurs ~10 μ s after the pulse start. In turn, the EMCCD camera trigs the high-speed camera with negligible delay (of <1 μ s) as the acquisition occurs, therefore one acquisition corresponds to one pulse.

High-speed imaging of the discharge is performed with rate of 100,000 fps and 128×96 pixels image resolution for current regimes of 450 and 650 A, and with rate of 75,000 fps and 128×128 pixels image resolution for current regimes of 800 and 1000 A. Exposure time for all experiments is 0.95 µs. Imagining is performed with 1/100 ND filter to avoid an overexposure.

Electric parameters of the experimental setup for all four current regimes are presented in Table 1. The values are averaged throughout all the measurements done within each current regime (~20–25 experiments at each current value).

I _{disch} , A ¹	U_c , V 2	P, kW	$\tau_{raise},\mu s^3$	$ au_{disch'}\mu s^{4}$	$\tau_{total}, \mu s$
450	120	54	90	230	320
660	150	100	95	240	335
800	180	146	100	240	340
1000	220	217	96	240	336

Table 1. Electric parameters of the experimental setup corresponding to different current regimes (per pulse).

Notes: ¹ I_{disch} corresponds to the peak value of the current; ² U_c corresponds to the load capacitor voltage; ³ τ_{raise} corresponds to the duration of the current raise up to its maximum value; ⁴ τ_{disch} corresponds to the duration of the current decrease.

2.2. Optical Emission Spectroscopy (OES)

2.2.1. Registered Spectra

The typical spectra of hydrogen Balmer and Cu I lines registered for different experimental regimes are shown in Figure 2. Figure 3 presents the spatially resolved spectrum images of the selected lines. It should be noted that no trace of hydroxyl radicals (OH) was recorded throughout the experiments. The possible explanation for this can be the fact that electrohydraulic spark and arc discharges are quite different from the "partial" discharges, i.e., streamer and corona discharges. The current between electrodes is transferred here by electrons. Because of the relatively high breakdown electric field of water, a small interelectrode gap is necessary and the discharge current heats a small volume of plasma, which results in the generation of almost thermal plasma (the temperatures of the electron and the heavy particles are almost equal, as known from measurements on a water-stabilized plasma
torch) [11]. Thermal plasmas pose a high risk of electrode erosion and are not very effective in the generation of radicals [18].

2.2.2. Broadening of Spectral Lines

The Stark broadening of the spectral lines is often used as a justified technique for the diagnosis of electron density (N_e), with applications not only in laboratory plasmas but also in astrophysical ones. This broadening is due to the collisions of the emitter with the charged particles in the surroundings of the emitter, according to the Stark effect. Stark broadening of the Balmer lines is the most popular approach to the determination of *Ne* since the broadening of the hydrogen lines, resulting from the linear Stark effect, is the strongest. These lines are, thus, those most sensitive to electron density variations. For the purposes of completeness and comparison, the electron density was also determined from the profile of Cu I 515.3 nm line, which is exposed to the quadratic Stark effect. In addition to Stark broadening, ideally, one should consider other broadening mechanisms, namely instrumental broadening, natural broadening, Doppler broadening, pressure broadening (resonance broadening, Van der Waals broadening), which have been reviewed in detail in [19,20]. More information about the broadening phenomena (physical phenomena, mathematical expressions of FWHMs, convolution procedure, etc.) are presented in Appendix A.



Figure 2. Typical spectra obtained throughout the experiments (cross-section of the arc halfway between the electrodes); exposure time of the EMCCD τ = 200 µs: (**a**) spectrum of Cu I lines registered for I = 450 A, central wavelength λ = 515.0 nm, grating 300 lines/mm; (**b**) H_{α} line registered for I = 450 A, central wavelength λ = 616.5 nm, grating 300 lines/mm; (**c**) H_{β} line registered for I = 660 A, central wavelength λ = 486.1 nm (H β line wasn't registered for the current regime of 450 A due to its low intensity), grating 1800 lines/mm.



Figure 3. Spatially resolved spectrum images of the selected lines; exposure time of the EMCCD τ = 200 µs: (a) spectrum of Cu I lines registered for I = 400 A; (b) H_{α} line registered for I = 400 A; (c) H_{β} line registered for I = 660 A.

2.2.3. Line Broadening Values in Our Configuration

In our work, apparatus function has a Gaussian shape with a widths of 0.012 nm for the 300 lines/mm grating and 0.001 nm for the 1800 lines/mm grating. The order of magnitude of obtained w_N values showed to be of $\sim 10^{-5}$ nm and is negligible in comparison to the impact of other broadening mechanisms. Values of w_R and w_{VdW} are of the order of magnitude of $\sim 10^{-3}$ nm and doesn't vary significantly with the values of gas temperature T_g , and as well can be excluded out of calculations of the electron density. Therefore, profiles of the registered lines are fitted with a Voigt function, whereas the Gaussian part is presented by convolution of the instrumental and Doppler broadening, giving the values of 0.02-0.04 nm, and the Lorentzian part of the profile is considered to be only the Stark FWHM.

3. Results

3.1. Electron Density

Line shapes and Voigt fit of the studied lines are shown in Figure 4. One can see in Figure 5 the radial profiles of electron number density for 1–2 mm diameter plasma. It can be noted that in Figure 4c there is a large uncertainty concerning the maximum of the line (between 875 a.u. and 975 a.u.) and, therefore, the FWHM. A higher maximum leads to a FWHM larger than 0.6 nm and therefore a larger electron density, but the Voigt fit was performed according to procedure recommended in [21] with the fixed Gaussian "contribution" into the line's width, which resulted in the profile shown in Figure 4c.



Figure 4. Line profiles and Voigt fit of the registered lines for current regime I = 1000 A, radial position r = 0 mm (discharge axis): (a) Cu I 515.3 nm line; (b) H_{α} line; (c) H_{β} line.



Figure 5. Radial profiles of the electron density calculated using broadening of the selected registered lines for different experimental current regimes: (**a**) I = 450 A, *Ne* calculated using Cu I 515.3 nm line and H_{α} line; (H_{β} line was not registered for this regime); (**b**) I = 660 A, *Ne* calculated using Cu I 515.3 nm, H_{α} and H_{β} lines; (**c**) I = 800 A, *Ne* calculated using Cu I 515.3 nm, H_{α} and H_{β} lines; (**d**) I = 1000 A, *Ne* calculated using Cu I 515.3 nm, H_{α} and H_{β} lines;

3.2. High-Speed Imaging

High-speed images of the discharge for different current regimes are presented in Figure 6, wherein the first image corresponds to a time about 10 µs after the pulse start. One can clearly see the bright spherical area corresponding to the discharge itself occurring into the bubble. Therefore, the information about the discharge and bubble development as a function of time and electrical parameters can be obtained. Figure 7 shows the images of evolution of bubble's size up to its maximal diameter for every current regime studied in this work. Dependence of the bubble's size on duration of the discharge for different current regimes is presented in Figure 8, and Figure 9 shows evolution of the bubble's size depending on a discharge current for different regimes. The size of the bubble was estimated from the correspondence of 1 mm to number of pixels obtained from the photograph of the ruler made for scale. Figure 9 presents two values for the same current, the first half of the curve (the "upper" part) corresponds to rising current and the second half (the "lower" part, on the right side) to the falling current. This allows to see the bubble's growth differs from its collapse.



Figure 6. High-speed image recording of the discharge for different current regimes: (**a**) I = 450 A, time interval between presented frames–60 μ s, image resolution–128 \times 96 px, frame rate–100,000 fps; (**b**) I = 660 A, time interval between presented frames–60 μ s; image resolution–128 \times 96 px, frame rate–100,000 fps; (**c**) I = 800 A, time interval between presented frames–65 μ s; image resolution–128 \times 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 \times 128 px, frame rate–75,000 fps.



Figure 7. High-speed imaging of the discharge and bubble size development up to the maximal size for different current regimes: (**a**) I = 450 A, time interval between presented frames–60 μ s, image resolution–128 × 96 px, frame rate–100,000 fps; (**b**) I = 660 A, time interval between presented frames–60 μ s; image resolution–128 × 96 px, frame rate–100,000 fps; (**c**) I = 800 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps; (**d**) I = 1000 A, time interval between presented frames–65 μ s; image resolution–128 × 128 px, frame rate–75,000 fps



Figure 8. Time evolution of a bubble's size for different values of maximal current.



Figure 9. Evolution of the bubble's size depending on a discharge current (its evolution in time within one pulse) for different values of maximal current: (**a**) for maximal current of 450 A; (**b**) for maximal current of 660 A; (**c**) for maximal current of 800 A; (**d**) for maximal current of 1000 A.

3.3. Erosion of Electrodes

Measurements of erosion were performed by controlling the weights of cathode and anode before and after the experiments using digital scales with precision of 0.1 mg. Table 2 presents the data on electrodes' erosion for current regime I = 450 A and I = 800 A, for comparison. Unfortunately, measurements of erosion were not carried out for current of 1 kA, since the electrode deformation after one pulse was so important that we needed to sharpen it again before the next pulse and the corresponding erosion was too low to get sufficient measurement accuracy.

Current	Number of	Erosion of	Erosion of	Total Erosion,	Erosion Per
(I, A)	Consecutive Pulses	Anode, g	Cathode, g	g	Pulse (g/Pulse)
450	15	0.0038	0.0005	0.0043	0.00028
800	15	0.004	0.0012	0.0052	0.00034

Table 2. Erosion of the copper electrodes after the carried out experiments.

4. Discussion and Conclusions

All the measurements and calculations performed within these experimental series allow us to make the following conclusion through the analysis of the results:

- For all the four current regimes studied in this work (I = 450 A, I = 660 A, I = 800 A and I = 1000 A), duration of the pulse in average shows to be 320–346 μ s with the rise time of a current up to its maximum value of 90–100 μ s (see Table 1). While raise time of a current up to its maximum value is the shortest for the current regime I = 450 A, the maximum bubble's size is reached the fastest for the case of the current regime of I = 800 A, for which the current raise time is the longest. The only pattern that can be distinguished (see Figures 7 and 9) is that the bubble reaches it maximum size not at the same time when the current value is on peak, but before that, maintaining its size more or less constant while current value equals to its maximum value for the given current regime. In order to be able to make any conclusions, more experiments must be performed, which will give the more reliable statistical data.
- The erosion does not change much for the different current regimes. Table 2 shows that for current of 800 A, which is approximately two times greater than the first current regime, increase of the eroded mass per pulse is ~20%. Erosion of anode shows to be greater than that of cathode (in our experiments anode is the moving electrode). This is consistent with the results obtained by other authors studying the submerged pulsed arc discharges [22], and may have one of the possible explanations that the energy dissipated in the anode is larger than in the cathode. As expected, the size of craters produced on the electrodes during arcing (Figure 10) increases with the measured erosion. The larger craters were formed on the anode where the erosion was larger.
- Calculation of the electron density shows (Figure 5) disagreement the values of *Ne* calculated using the Cu I 515.3 nm, H_{α} and H_{β} lines. While *Ne* determined from the widths of Cu I 515.3 and H_{α} line are of the same order of magnitude for all the studied current regimes, values of *Ne* obtained from the width of H_{β} line are by two orders of magnitude lower for cases of I = 660 and 800 A, and by one order of magnitude lower for the case of I = 1000 A.

Normally, the use of the H_{α} line can lead to the following problems:

- It could present a non-negligible self-absorption and it is necessary to evaluate how it affects the line broadening;
- It has a strong broadening by ion dynamics, an effect that can be evaluated by using some computational methods recently developed which allow us to simulate the profiles of the spectral lines [23].



Figure 10. Photographs of the cathode (**left**) and anode (**right**) after the series of 15 experiments for I = 450 A. Diameter of both electrodes is 6 mm.

Taking into account the fact that self-absorption effects were not considered in the present study, one can conclude that the inconsistency between the values of *Ne* is caused by overestimation of the H_{α} line width. However, at the same time, values of *Ne* obtained from H_{α} line and Cu I 515.3 nm line show good agreement leading to the conclusion that the suggested value of the electron temperature Te = 10,000 K and $N_e \sim 10^{23}$ m⁻³ are close to the experimental values of the electron density and electron temperature in our discharge.

It was shown in [24] that the expression connecting the Stark width of Balmer lines with N_e, which has been derived from the Kepple–Griem theory (KG) [25], in case of use of H_{α} line, overestimates the electron density by about 80% with respect to the *Ne* obtained from the H_{β} line. This overestimation is due to the KG theory not taking into account the influence of the ion dynamics on the spectral profiles of the Balmer series lines. Computational methods taking into consideration these ion dynamic effects have been developed, among them the Gigosos–Cardenoso model (GC) [26] permitting the use of a line different from H_{β} for measuring the electron density, obtaining in this way the same value of density as the one obtained from H_{β} .

Values of the FWHM of H_{α} and H_{β} lines with the corresponding *Ne* and *Te* = 10,000 K for the case of μ = 0.9 (reduced perturber mass corresponding to the hydrogen emitter and any perturbing atom or ion) are showed in Table 3.

$\log N_e (m^{-3})$	FWHM (nm) for $H_{\boldsymbol{\alpha}}$	FWHM (nm) for $H_{\boldsymbol{\beta}}$
20.00	0.0142	0.0424
20.33	0.0244	0.0741
20.67	0.0404	0.129
21.00	0.064	0.217
21.33	0.102	0.361
21.67	0.16	0.601
22.00	0.25	0.999
22.33	0.393	1.67
22.67	0.621	2.80
23.00	1.01	4.70
23.33	1.68	7.77
23.67	2.84	1.27
24.00	4.86	2.05
24.33	8.31	3.25

Table 3. FWHM (in nm) of H_{α} and H_{β} lines for $\mu = 0.9$ (for Te = 10,000 K) [23].

It can be seen from the Table 3 that in order to get the same *Ne* at *Te* = 10,000 K, FWHM of H_{β} line must be between 3 and 4 times greater than that of H_{α} , while the experimentally obtained values of FWHM of H_{β} line are around 2–6 times smaller than that of H_{α} . (see Table 4), which, to our beliefs, leads to the correspondingly lower values of the estimated N_e.

Table 4. Experimentally obtained values of FWHM of H_{α} and H_{β} lines (in nm).

Line	FWHM (nm) at I = 660 A	FWHM (nm) at I = 800 A	FWHM (nm) at I = 1000 A	Average log N_e (m ⁻³)
Hα	1.13	2.60	2.40	23.36
H_{β}	0.40	0.39	0.57	21.51

It can be seen from Tables 3 and 4 that the obtained results of FWHM of H_{α} line are in the 30–33% agreement with the data given by GC in [23] in terms of $N_e/FWHM$ ratio.

Therefore, the conclusion can be made that it is important to estimate the self-absorption of the H_{α} line in order to double-check the results. Moreover, other techniques for the electron density determination should be used to validate the obtained results, for instance, determination of the electron density from continuum radiation, or, for instance, technique proposed in [27].

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Appendix A.

Whenever the contribution of other broadening mechanisms compared to Stark broadening is not negligible, the deconvolution procedure must be employed to determine Stark width– w_s , which is used to determine *Ne* from the comparison with theoretical widths at the same electron temperature (T_e).

One should consider the following mechanisms contributing to the spectral line profile and width:

- Instrumental broadening;
- Natural broadening;
- Doppler broadening;
- Pressure broadening (resonance broadening, Van der Waals broadening).

The deconvolution of the various contributions in the line broadened profile can be carried out based on the fact that, in many cases, the experimentally measured line profiles can be fitted with the Voigt function which is the convolution of the two functions: Gaussian and Lorentzian with the full width at half maximum (FWHM) given by w_G and w_L . The FWHM of a Voigt profile can be represented through the combination of the Gauss and Lorentz components as follows [21]:

$$w_v = \frac{w_L}{2} + \left(\left(w_L/2\right)^2 + w_G^2\right)^{1/2}.$$
 (A1)

The broadening contributions with a Gaussian shape will lead to a profile with w_G determined as:

$$w_G = \left(\sum_i (w_{Gi})^2\right)^{1/2}.$$
 (A2)

Whereas for Lorentzian broadening components the final contribution to the line profile will also be Lorentzian, with FWHM determined as:

$$w_L = \sum_i w_{Li}.$$
 (A3)

Appendix A.1. Instrumental Broadening

Instrumental broadening is important, in particular at low *Ne* and low emitter's temperature when line profile is narrow, i.e., when Stark broadening is small or comparable with other broadening mechanisms. The standard practice to determine the apparatus profile of a spectrometer is to scan over a line whose intrinsic width is very small compared to the apparatus width so that the latter determines its shape. Typically, such narrow lines are obtained from low-pressure discharges, like Geissler tubes or hollow cathode lamps. The instrumental line shapes obtained with good spectrometers are usually approximately Gaussian profiles in particular central core which usually is being used for deconvolution procedure.

Appendix A.2. Natural Broadening

Natural broadening is a consequence of the Heisenberg uncertainty principle applied to the energies of the initial and the final states of the transition. The uncertainty on the transition frequency or wavelength translates into a Lorentzian line profile of FWHM, for transition between states n and m [19]:

$$w_N(\mathrm{cm}) = \frac{\lambda_{nm}^2 (\sum_{u < n} A_{nu} + \sum_{u < m} A_{mu})}{2\pi c},\tag{A4}$$

where λ_{nm} is the wavelength of a spectral line corresponding to transition from an upper state *n* to the lower state *m* (in cm), A_{nu} , A_{mu} are the transition probabilities of transitions between the states *n*, *m* and any 'allowed' level *u*, correspondingly Natural broadening is largest when one of the two levels is dipole-coupled to the ground state. Even in this case, it is usually negligible (of the order of 10^{-4} nm). However, it may be of some importance for low electron density plasmas generated in low-pressure gas discharges. The values of the Einstein *A* coefficients for calculations were taken from [28].

Appendix A.3. Resonance Broadening

Resonance broadening occurs for transitions involving a level that is dipole-coupled to the ground state and can be derived from the following equation [19]:

$$w_{res} (cm) = 1.63 \times 10^{-13} (g_i / g_k)^{1/2} \lambda^2 \lambda_R f_R N_a,$$
(A5)

where λ is the wavelength (in cm) of the observed radiation. Here, N_a is the density of ground state particles which are the same species as the emitter, and g_i and g_k are the statistical weights of the upper and lower state; λ_R and f_R are the wavelength and f-value (oscillator strength), respectively, of the resonance transition from level "R". The level "R" is the upper or lower level of the observed transition, which happens also to be the upper level of a resonance transition to the ground state. Not all transitions involve such a level. The values of f_R , g_i and g_k for calculations were taken from [28].

Appendix A.4. Van der Waals Broadening

Van der Waals broadening results from the dipole interaction of an excited atom with the induced dipole of a neutral ground-state atom of density N_a . This is a short-range C_6/r_6 interaction. The full-width at half-maximum of the Lorentzian profile w_{VdW} can be written as [19]:

$$w_{VdW}$$
 (cm) = $8.18 \times 10^{-12} \lambda^2 \left(\overline{\alpha} \overline{R^2}\right)^{2/5} (T_g/\mu)^{3/10} N_a$, (A6)

where

$$\overline{R^2} = \overline{R_U^2} - \overline{R_L^2} \tag{A7}$$

 $\overline{R^2}$ is the difference of the squares of coordinate vectors (in a_0 units) of the upper and lower level, and μ is the atom-perturber reduced mass in a.m.u. In the Coulomb approximation the values of $\overline{R_U^2}$ and $\overline{R_L^2}$ in (7) may be calculated from Atoms 2017, 5, 40

$$\overline{R_j^2} = n_j^{*2} / 2 \Big[5n_j^{*2} + 1 - 3l_j (l_j + 1) \Big],$$
(A8)

where the square of effective quantum number n_i^{*2} is

$$n_{j}^{*2} = E_{H} / \left(E_{IP} - E_{j} \right) \tag{A9}$$

where E_H is the ionization potential of hydrogen (109,737.32 cm⁻¹). Here E_{IP} is the ionization potential of the studied element and E_i is the energy of the upper or lower level of the transition.

Values of $\overline{\alpha}$, the mean atomic polarizability of the neutral perturber, are tabulated for different elements in [29]. If the required value of $\overline{\alpha}$ is not tabulated, it can also be estimated from the expression given in [29]:

$$\overline{\alpha} = (9/2)a_0^3 (3E_H/4E_{EXC})^2, \tag{A10}$$

where E_{EXC} is the energy of the first excited level of the perturber.

Van der Waals broadening causes a red wavelength shift which is two-third of the size of the width w_{VdW} . In our case, we consider that discharge occurs at atmospheric pressure (or higher) in air with 1–2% of the hydrogen, similar to [30]. Therefore, value of $\overline{\alpha}$ presents a sum of atomic polarizabilities of air components (O, N, O₂, N₂, NO), and it was not taken from [29], but was calculated using Equation (A8). Mean atom-perturber reduced mass μ is taken to be equal to one (reduced perturber mass corresponding to the hydrogen emitter and any perturbing atom or ion).

Appendix A.5. Doppler Broadening

The relative motion of an emitter to the detector leads to a shifted line. In plasma with gas temperature T_g the line profile due to the presence of Doppler effects can be well described by a Gaussian profile with a FWHM (nm):

$$w_D = 1/2\lambda_0 \sqrt{\frac{8kT_g ln2}{m_a c^2}},\tag{A11}$$

where *k* is the Boltzmann constant (unit: J K⁻¹) and T_g the gas temperature in *K*, and m_a –the mass of the emitter (a.m.u.).

Appendix A.6. Stark Broadening

The analysis of Stark-broadened spectral-line profiles is one of the most common methods of plasma diagnosis from the beginning of the sixties. The shape of these profiles depends heavily on the density of charged particles surrounding the emitter. This dependence is especially important in hydrogen and hydrogenic ions, since in those elements a linear Stark effect takes place. As a consequence, Stark-broadening measurements have become one of the most widely used techniques when plasma electron density is to be determined. On the other hand, accurate knowledge of the shapes of the line profiles are mandatory in order to determine the opacity spectral distribution. These kind of diagnostics play a key role in astrophysics, where active spectroscopy is not possible.

To this end, profile tables of hydrogen Stark broadening have been in use since the beginning of the seventies [20,31]. The calculations leading to these tables were made in the framework of a plasma model in which it is assumed that the broadening of the lines can be computed as a sum of the contributions of all the collisions of a statistical ensemble of quasi-independent charged particles with the emitting atom. Depending on the speed of the perturbing particles, two different approximations were considered in order to simplify the computation of the contribution of each of the two different broadening mechanisms which are taken into account: it is assumed that the highly mobile electrons are responsible for sudden variations in the phase of the emitted wave–impact regime, while the

heavier and slower ions generate electric fields which can be considered constant along the life of a typical emission–quasistatic regime.

The admissibility of this approximation seemed to be well secured for typical laboratory plasmas [20], and for that reason theoretical research in the field has concentrated mainly on electron broadening. The greater was the surprise when Kelleher and Wiese [32] and Wiese et al. [33] proved experimentally that the central parts of the first Balmer lines depend markedly on the reduced mass of the radiating atom-perturbing ion pair.

While these experimental results were obtained, other Stark-broadening calculation methods were developed [26,34].

Both the linear and the quadratic Stark effects are encountered in spectroscopy. However, only the hydrogen atom and H-like ions exhibit the linear Stark effect, whereas all other atoms exhibit the quadratic Stark effect. In the case of linear Stark effect, the relation between electron density and the line width can be expressed as a simple relation (static ion approximation) [35], being extremely convenient for the estimation of order of magnitude of the electron density value:

$$N_e = C(N_e, T_e) \times (w_S)^{3/2},$$
(A12)

where w_S is the FWHM of a Lorentzian contribution to the overall line profile and the parameter $C(N_e, T_e)$ depends (only weakly) on N_e and T_e , which can normally be treated as being constant. The constant *C* for *H* Balmer lines is available in the literature [35]. Usually, the first choice for the electron density determination is H_β line (with an error of 5%) [35] because of its large intensity and sufficiently large line broadening, which can be measured precisely using spectrometer of moderate resolution. The possibility of self-absorption in this case is relatively small. The second best choice among the Balmer series is the H_v line. The H_α line is suitable in the cases where the electron density is not too high ($N_e \sim 10^{17} \text{ cm}^{-3}$), because at higher electron densities this strong line is quite susceptible to self-absorption, which severely distorts the line profile.

In the case of non-H-like atoms, where the quadratic Stark effect is dominant, the relation between the electron density and the line width is [35]:

$$w_S \approx 2[2 + 1.75 \times 10^{-4} N_e^{\frac{1}{4}} \alpha (1 - 0.068 N_e^{\frac{1}{6}} T^{-1/2})] \times 10^{-16} w N_e.$$
(A13)

The first term in the brackets gives the contribution from the electron broadening, and the second term stems from ion broadening. Here w is the electron impact parameter at $N_e = 10^{16}$ cm⁻³, and α is the ion broadening parameter. These parameters can be found easily from the literature [35]. Since the second term after opening the brackets in Equation (A13) is normally small, so the expression reduces to

$$w_S \approx 4 \times 10^{-16} w N_e. \tag{A14}$$

It has been reported in [36] that the electron density derived from the H_{β} line and H_{v} line data were in good agreement with each other, while the electron density estimated from the H_{α} line is not purely Stark broadened. The additional broadening might be due to the onset of self-absorption.

In this work, values of electron density were obtained using H_{β} , H_{α} lines from the following equations presented in [26], which takes into account the ion dynamic effects:

for
$$H_{\beta}$$
: $N_{e} \left[m^{-3} \right] = 10^{23} \times (w_{S} [nm]/4.8)^{1.46808}$, (A15)

for
$$H_{\alpha}: N_{e}\left[m^{-3}\right] = 10^{23} \times (w_{S}[nm]/1.098)^{1.47135}$$
. (A16)

Electron density for Cu I 515.3 nm line is obtained from the linear interpolation from the table presented in [37], whereas the theoretical and experimental Stark widths of Cu I lines are listed for

 $N_e = 10^{17}$ cm⁻³ and $T_e = 10,000$ K. These values were assumed as parameters for the estimations presented in this work.

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