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Complex many-electron highly charged ions studies at ITPA, Vilnius, Lithuania

Monday, October 5, 2009
About 3,500,000 inhabitants live in Lithuania.
VILNIUS is the capital of the Republic of Lithuania. The population of Vilnius is more than 542,000. Current area of Vilnius is 392 square kilometres. The Old Town, historical centre of Vilnius, is one of the largest in Eastern Europe (360 ha). The most valuable historic and cultural heritage is concentrated here. Because of its uniqueness, the Old Town of Vilnius was inscribed on the UNESCO World Heritage List.
Institute of Theoretical Physics and Astronomy of Vilnius University

It is a budgetary State research body, entitled to perform research in theoretical physics and astronomy, and aiding the Vilnius University and other higher education establishments in preparing the scientists and specialists of high qualification.

It was established in 1990 on the basis of 5 departments which had previously belonged to the Institute of Physics in Vilnius though the departments belonging to it appeared and were continuously developing since 1952 at the institute of the Academy of Sciences that had been existing under different names.

Since 2002 the Government of the Republic of Lithuania has granted the university research institute status to the ITPA.
The following main research directions are:

• Development of effective methods of mathematical physics and their application in theoretical investigation of many-particle systems, their non linear dynamics and of quantum fields.

• Investigations of atoms, subatomic particles, molecules, their structures and plasma spectroscopy, their application in nanophysics and astrophysics.

• Investigations of the structure and evolution of the Galaxy, stars and interstellar matter.
Structure

- Astronomical Observatory (former Department of Astrophysics)
- Department of the Theory of an Atom
- Department of Processes and Structures
- Department of Isotopic Analysis
- Department of Nucleus of Atom
- Planetarium
Astronomical Observatory

The main research areas:

• Multicolor stellar photometry,
• Stellar classification, chemical analysis of stellar atmospheres,
• Interstellar extinction,
• Galactic structure and chemical evolution,
• Stellar radial velocities,
• The search and positional observations of comets, asteroids and near-Earth objects.
Astronomical Observatory

Astronomers of ITPA AO participate in a number of international projects, including photometric investigation of Galactic areas (with USA and Vatican astronomers), spectral investigations of evolved stars (with astronomers from Germany, Italy, Sweden, Switzerland, USA, Canada and Chile), investigations of galaxy evolution (with astronomers from England), photometric, spectral and dynamical investigation of open clusters (with Taiwan astronomers), photometric investigation of open clusters in the Magellanic Cloud galaxies (with astronomers of Japan), the Whole Earth Telescope program for investigation of variable white dwarfs, the Gaia orbiting observatory project of ESA, the Kepler orbiting observatory project of NASA.
The construction of the Moletai Astronomical Observatory was started in 1964. It is situated about 70 km from Vilnius and owns 165 cm and 63 cm reflecting telescopes and the 35/51 cm Maksutov-type telescope. The main instruments of astronomical observations are a CCD photometer, three-channel and two-channel photoelectric photometers and a Coravel-type instrument for radial velocity measurements.
Planetarium founded in 1962. There are 150 places in its lecture hall. The lectures about astronomy, physics, geography and nature are given to pupils, students and broad public.
Department of the Theory of an Atom (ATD)

The main research directions are:

• Theoretical atomic spectroscopy,
• The development of methods of the theory of complex atomic and ionic spectra,
• The development of quantum many body theory,
• The algorithms and computer programs are worked out and applied in plasma physics, astrophysics and other fields.
Fairly large group of theoretical physicists of the Research Institute of Theoretical Physics and Astronomy of Vilnius University is involved in the development of the theory of complex many-electron atoms and ions, highly ionized atoms included, and various applications of the theory.

A summary of the topics is as follows:

1. Theoretical atomic spectroscopy, methods of the theory of complex atomic and ionic spectra, symmetry properties of the systems of the considerations;
2. Algorithms and programs for the calculation of energy spectra, radiative and autoionization probabilities and radiative lifetimes, taking into account relativistic and correlation effects;

3. Sophisticated large scale calculations of wavelengths as well as electric and magnetic multipole transition probabilities for many-electron atoms and ions;

4. Investigation of processes in the inner shells of atoms, x-ray and Auger transitions as well as their cascades;

5. Application of the global characteristics method for the approximate description of complex atomic spectra and investigation of regularities in these spectra;
6. Application of atomic theory for the description of the polarization in the interaction of atoms and ions with photons, electrons and other charged particles;

7. Investigation of polarization patterns of radiation and Auger electrons following excitation and ionization of atoms and ions by electron and photon impact;

8. Simulation of the radiation spectrum of impurities in tokamak plasma, applications to laser produced plasma and astrophysics.
ATD Staff working in atomic theory

Prof. **Zenonas Rokus Rudzikas** – Head
Prof. **Romualdas Karazija** – Chief researcher
Prof. **Pavel Bogdanovich** – Chief researcher
Prof. **Gediminas Gaigalas** – Chief researcher
Dr. **Vladas Tutlys** – Senior researcher
Dr. **Alicija Kupliauskienė** – Senior researcher
Dr. **Sigitas Kučas** – Senior researcher
Dr. **Gintaras Merkelis** – Senior researcher
Dr. **Romualdas Kisielius** – Senior researcher
Dr. **Valdas Jonauskas** – Senior researcher
Dr. **Rasa Karpuškienė** – Senior researcher
Dr. **Alina Momkauskaitė** – Researcher
Dr. **Aušra Kynienė** – Researcher
Dr. **Olga Rancova** – Researcher

**Rytis Juršėnas** – PhD student
Prof. Adolfas Jucys (1904 – 1974)
Prof. Adolfas Jucys is the founder of the school of theoretical physics in Lithuania. His most important results are:

The general form of the multiconfiguration Hartree-Fock equations (Hartree-Fock-Jucys equations) was derived for the first time.


A graphical technique of the angular momentum


ADOLFAS JUCYS

Teorinis ionų C^4+ ir C^{++}
ir Neutralaus C Tyrimas

Theoretical Investigation of Ions C^4+ and C^{++},
and of Neutral C

Disertacija daktaro laipsniui gauti Vilniaus
Universiteto Matematikos-Gamtos Fakultete,
A. P. YUTSIS
I. B. LEVINSON
V. V. VANAGAS

THEORY OF
ANGULAR MOMENTUM

Translated from Russian

Published for the National Science Foundation, Washington D. C.
and the National Aeronautics and Space Administration by the
Israel Program for Scientific Translations
Development of the theory

- Applications of the second quantization methods
- Transformed radial wave functions in the configuration interaction method (CI)
- A creation of new computer algorithms
- Applications of general spectral characteristics for complex spectra
- Applications of the methods of the theory of an atom for the investigation of polarization in the interaction of atoms with photons and electrons.
In quantum mechanics, the physical quantities are proportional to the squares of matrix elements.

The matrix element of the physical operator can be written as the product of the angular and radial parts.

For each of these parts, the distinctive methods are used because of specific problems in their evaluation.
Theoretical Atomic Spectroscopy

ZENONAS RUDZIKAS

CAMBRIDGE MONOGRAPHS ON ATOMIC, MOLECULAR AND CHEMICAL PHYSICS
Angular coefficients

A general method for finding algebraic expressions for matrix elements of one- and two-electron operator for an arbitrary number of subshells in an atomic configuration requiring neither coefficients of fractional parentage nor unit tensors is developed by applying quasispin and isospin quantities.

These expressions are implemented into computer programs and up to hundreds of times shorten the calculations.

Any two-particle operator

\[
\tilde{G}^{(\kappa_1 \kappa_2 k, \sigma_1 \sigma_2 k)} = \sum_{\alpha} \sum_{\kappa_1 \kappa_2 k, \sigma_1 \sigma_2} \Theta(\Xi) \left\{ A^{(k \ k)}_{p, -p} (n_\alpha \lambda_\alpha, \Xi) \delta(u, 1) \right. \\
+ \sum_{\beta} \left[ B^{(\kappa_{12} \sigma_{12})} (n_\alpha \lambda_\alpha, \Xi) \times C^{(\kappa'_{12} \sigma'_{12})} (n_\beta \lambda_\beta, \Xi) \right]^{(k \ k)}_{p, -p} \delta(u, 2) \\
+ \sum_{\beta \gamma} \left[ D^{(l_\alpha s)} \times D^{(l_\beta s)} \right]^{(k \ k)}_{\kappa_{12} \sigma_{12}} \times E^{(\kappa'_{12} \sigma'_{12})} (n_\gamma \lambda_\gamma, \Xi) \left]^{(k \ k)}_{p, -p} \delta(u, 3) \\
+ \left. \sum_{\beta \gamma \delta} \left[ D^{(l_\alpha s)} \times D^{(l_\beta s)} \right]^{(k \ k)}_{\kappa_{12} \sigma_{12}} \times \left[ D^{(l_\gamma s)} \times D^{(l_\delta s)} \right]^{(k \ k)}_{\kappa'_{12} \sigma'_{12}} \right \} \left]^{(k \ k)}_{p, -p} \delta(u, 4) .
\]

The tensorial part of a two–particle operator is expressed in terms of operators of the type \( A^{(k \ k)}_{p, -p} (n_\alpha \lambda_\alpha, \Xi) \), \( B^{(\kappa_{12} \sigma_{12})} (n_\alpha \lambda_\alpha, \Xi) \), \( C^{(\kappa'_{12} \sigma'_{12})} (n_\beta \lambda_\beta, \Xi) \), \( D^{(l_\alpha s)} \) and \( E^{(\kappa'_{12} \sigma'_{12})} (n_\gamma \lambda_\gamma, \Xi) \).

The amplitude \( \Theta(\Xi) \) is proportional to the two–electron submatrix element \( (n_i \lambda_i \ n_j \lambda_j || g || n_{i'} \lambda_{i'} \ n_{j'} \lambda_{j'}) \).
Matrix Elements Between Complex Configurations

\[
G \left| \langle \psi_u (LS) || \psi_u (L'S') \rangle \right|
\]

\[
= \sum_{n_i l_i, n_j l_j, n'_i l'_i, n'_j l'_j} \langle \psi_u (LS) || \hat{G} \left( n_i l_i, n_j l_j, n'_i l'_i, n'_j l'_j \right) || \psi_u (L'S') \rangle
\]

\[
= \sum_{n_i l_i, n_j l_j, n'_i l'_i, n'_j l'_j} \sum_{\kappa_{12}, \sigma_{12}, \kappa'_{12}, \sigma'_{12}} \sum_{K_l, K_s} (-1)^{\Delta \Theta'} \left( n_i \lambda_i, n_j \lambda_j, n'_i \lambda'_i, n'_j \lambda'_j, \Xi \right)
\times T \left( n_i \lambda_i, n_j \lambda_j, n'_i \lambda'_i, n'_j \lambda'_j, \Lambda_{bra}^bra, \Lambda_{ket}^ket, \Xi, \Gamma \right)
\times R \left( \lambda_i, \lambda_j, \lambda'_i, \lambda'_j, \Lambda_{bra}^bra, \Lambda_{ket}^ket, \Gamma \right),
\]

where \( \lambda \equiv l, s, \Lambda_{bra}^bra \equiv \left( L_i, L_j, L'_i, L'_j \right)^{bra}, \Lambda_{bra}^bra \equiv \left( S_i, S_j, S'_i, S'_j \right)^{bra} \) and \( \Gamma \) refers to the array of coupling parameters connecting the recoupling matrix \( R \left( \lambda_i, \lambda_j, \lambda'_i, \lambda'_j, \Lambda_{bra}^bra, \Lambda_{ket}^ket, \Gamma \right) \) to the submatrix element \( T \left( n_i \lambda_i, n_j \lambda_j, n'_i \lambda'_i, n'_j \lambda'_j, \Lambda_{bra}^bra, \Lambda_{ket}^ket, \Xi, \Gamma \right) \). The expression has summation over intermediate ranks \( \kappa_{12}, \sigma_{12}, \kappa'_{12}, \sigma'_{12}, K_l, K_s \) in \( T \left( n_i \lambda_i, n_j \lambda_j, n'_i \lambda'_i, n'_j \lambda'_j, \Lambda_{bra}^bra, \Lambda_{ket}^ket, \Xi, \Gamma \right) \).
Spin-Angular Part of a Two Particle Operator

The tensorial part of a two–particle operator is expressed in terms of operators of the type $A^{(kk)}(n\lambda,\Xi)$, $B^{(kk)}(n\lambda,\Xi)$, $C^{(kk)}(n\lambda,\Xi)$, $D^{(ls)}$, $E^{(kk)}(n\lambda,\Xi)$. Their explicit expressions are:

\[ a^{(q\lambda)}_{m_q}, \]

\[ \left[ a^{(q\lambda)}_{m_{q1}} \times a^{(q\lambda)}_{m_{q2}} \right]^{(k_1\sigma_1)} \]

\[ \left[ a^{(q\lambda)}_{m_{q1}} \times \left[ a^{(q\lambda)}_{m_{q2}} \times a^{(q\lambda)}_{m_{q3}} \right]^{(k_1\sigma_1)} \right]^{(k_2\sigma_2)}, \]

\[ \left[ \left[ a^{(q\lambda)}_{m_{q1}} \times a^{(q\lambda)}_{m_{q2}} \right]^{(k_1\sigma_1)} \times a^{(q\lambda)}_{m_{q3}} \right]^{(k_2\sigma_2)}, \]

\[ \left[ a^{(q\lambda)}_{m_{q1}} \times a^{(q\lambda)}_{m_{q2}} \right]^{(k_1\sigma_1)} \times \left[ a^{(q\lambda)}_{m_{q3}} \times a^{(q\lambda)}_{m_{q4}} \right]^{(k_2\sigma_2)} \right]^{(kk)} \]

We denote their submatrix elements by $T \left( n_i \lambda_i, n_j \lambda_j, n_i' \lambda_i', n_j' \lambda_j', \Lambda^{bra}, \Lambda^{ket}, \Xi, \Gamma \right)$. 
The Quasispin Formalism

We obtain the submatrix elements of operator $a_{m_q}^{(q\lambda)}$ by using straightforwardly the Wigner–Eckart theorem in quasispin space:

$$
\left\langle l^N \alpha QLS | a_{m_q}^{(q\lambda)} | l'^{N'} \alpha' Q'L'S' \right\rangle
$$

$$
= - [Q]^{-1/2} \begin{bmatrix}
\frac{Q'}{M'_Q} & \frac{1/2}{m_q} & \frac{Q}{M_Q} \\
M'_Q & m_q & M_Q
\end{bmatrix} \left( l \alpha QLS || a_{m_q}^{(q\lambda)} || l' \alpha' Q'L'S' \right),
$$

where the last multiplier is the so-called reduced coefficient of fractional parentage. All standard quantities, which are considered in this approach, can be defined in terms of these coefficients.

The relation between coefficient of fractional parentage and the reduced coefficients of fractional parentage is:

$$
\left\langle j^N \alpha QJ | j^{N-1} (\alpha' Q' J') j \right\rangle
$$

$$
= \frac{(-1)^{N+1}}{\sqrt{N} [J, Q]} \begin{bmatrix}
\frac{Q'}{M'_Q} & \frac{1/2}{1/2} & \frac{Q}{M_Q} \\
M'_Q & 1/2 & M_Q
\end{bmatrix} \left( j \alpha QJ || a_{m_q}^{(q\lambda)} || j' \alpha' Q' J' \right).
Multiconfiguration wave function

\[ \langle K_1 \lambda LS \rangle = \sum_{K'T'} a(K_1 \lambda LS, K'T' LS) \langle K'T' LS \rangle. \]

For its determination two methods are used: **Configuration interaction** (CI) where the expansion coefficients are varied but the basis of radial wave functions is fixed, and **Multiconfiguration** where both expansion coefficients and radial orbitals are varied.
He $1s^2 + 2p^2$

Transformed radial orbitals

\[ P_{TRO} \{ \ell \mid r \} = N f(k, m, B \mid r) P_{HF} \{ \ell_0 \mid r \} \]

\[ f(k, m, B) = r^k \exp(-Br^m), \]
\[ k \geq l - l_0, \quad k > 0, \quad m > 0, \quad B > 0. \]

Transformed radial orbitals

• The basis of TRO can be generated easy and quickly.

• Thus, hundreds of thousands or even millions of configuration states can be taken into account for the investigation of correlation effects.

• But our computers can not get over such large matrices of the energy.

• To overcome this problem some improvements of the calculation algorithms were made.
Improvement of calculation algorithms

- **Selection of configurations** by using the second order of perturbation theory.
- **Special coupling scheme** of the shells of admixed configurations. That allows us to reduce the number of configuration states several times.
- **The partial diagonalization of matrices** by using the Jacobi method for the determination of the eigenvalues and eigenfunctions that are of interest.
- **The sequential diagonalization of the matrices**. At first all the necessary eigenvalues and then the corresponding eigenfunctions are determined. This method allows one to promptly diagonalize the symmetric matrices if only their triangular part fits into the computer RAM.
Improvement of calculation algorithms

The separate diagonalization of the energy matrices. At first all the energy matrices for the separate pairs of $LS$ of the operator $H$ are diagonalized. The interactions depending on the final momentum $J$ are included only for the adjusted and energetically close admixed configurations.

As the result of these operations the orders of the matrices for the separate $J$ values are small.

All these means enable one to reduce the order of the energy matrix from millions down to hundreds of thousands.
<table>
<thead>
<tr>
<th></th>
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<tbody>
<tr>
<td>3s²</td>
<td>1S₀</td>
<td>0</td>
<td>5</td>
</tr>
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<td>3s3p</td>
<td>3P₀</td>
<td>143176</td>
<td>142987</td>
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<td></td>
<td>3P₁</td>
<td>144675</td>
<td>144557</td>
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<td></td>
<td>3P₂</td>
<td>147912</td>
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<td>1P₁</td>
<td>214482</td>
<td>214482</td>
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<tr>
<td>3p²</td>
<td>1D₂</td>
<td>336245</td>
<td>336762</td>
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<tr>
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<td>3P₀</td>
<td>339963</td>
<td>339754</td>
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<td>3P₁</td>
<td>341872</td>
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<td></td>
<td>3P₂</td>
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<td>398776</td>
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<td>3D₂</td>
<td>412191</td>
<td>412582</td>
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<td></td>
<td>3D₃</td>
<td>412405</td>
<td>412946</td>
</tr>
<tr>
<td></td>
<td>1D₂</td>
<td>467631</td>
<td>468131</td>
</tr>
</tbody>
</table>

Oscillator strengths for the transitions $3p4p$-$3p3d$ for Ca IX

<table>
<thead>
<tr>
<th></th>
<th>$\text{L}$</th>
<th>$\text{V}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$1^1D_2$</td>
<td>$^3F_2$</td>
<td>$1.50 \times 10^{-3}$</td>
</tr>
<tr>
<td>$1^1D_2$</td>
<td>$^3F_3$</td>
<td>$7.70 \times 10^{-4}$</td>
</tr>
<tr>
<td>$1^1D_2$</td>
<td>$1^1D_2$</td>
<td>$6.93 \times 10^{-2}$</td>
</tr>
<tr>
<td>$^3P_2$</td>
<td>$^3F_3$</td>
<td>$3.36 \times 10^{-3}$</td>
</tr>
<tr>
<td>$^3P_1$</td>
<td>$^3F_2$</td>
<td>$1.21 \times 10^{-3}$</td>
</tr>
<tr>
<td>$^3D_3$</td>
<td>$^3F_2$</td>
<td>$1.35 \times 10^{-3}$</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>$^3P_2$</td>
<td>4.2±0.7</td>
<td>3.36±0.16</td>
<td>3.31</td>
<td>3.34</td>
</tr>
<tr>
<td>$^3F_3$</td>
<td>4.1±0.5</td>
<td>3.29±0.09</td>
<td>3.18</td>
<td>3.33</td>
</tr>
<tr>
<td>$^3P_1$</td>
<td>3.4±0.6</td>
<td>2.90±0.14</td>
<td>2.59</td>
<td>2.41</td>
</tr>
<tr>
<td>$^3F_4$</td>
<td>3.3±0.4</td>
<td>2.56±0.08</td>
<td>2.72</td>
<td>3.13</td>
</tr>
<tr>
<td>$^3D_2$</td>
<td>3.7±0.6</td>
<td>2.80±0.10</td>
<td>2.79</td>
<td>3.09</td>
</tr>
<tr>
<td>$^3P_0$</td>
<td>2.6±0.6</td>
<td>3.38±0.16</td>
<td>3.11</td>
<td>3.29</td>
</tr>
<tr>
<td>$^3D_3$</td>
<td>3.4±0.6</td>
<td>2.36±0.07</td>
<td>2.46</td>
<td>2.88</td>
</tr>
<tr>
<td>$^3F_2$</td>
<td>3.8±0.6</td>
<td>3.12±0.10</td>
<td>2.97</td>
<td>3.19</td>
</tr>
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</table>

## Radiative lifetimes (ns) of Ca IX

<table>
<thead>
<tr>
<th></th>
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<th></th>
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</thead>
<tbody>
<tr>
<td>3s3p</td>
<td>$^3$P$_2$</td>
<td>0.0940</td>
<td>0.0982</td>
<td>0.095±0.010</td>
</tr>
<tr>
<td>3p$^2$</td>
<td>$^1$D$_2$</td>
<td>0.872</td>
<td>0.907</td>
<td>0.84±0.060</td>
</tr>
<tr>
<td></td>
<td>$^3$P$_0$</td>
<td>0.112</td>
<td>0.118</td>
<td>0.134±0.008</td>
</tr>
<tr>
<td></td>
<td>$^3$P$_2$</td>
<td>0.110</td>
<td>0.113</td>
<td>0.150±0.008</td>
</tr>
<tr>
<td>3s3d</td>
<td>$^3$D$_1$</td>
<td>0.0765</td>
<td>0.0783</td>
<td>0.100±0.015</td>
</tr>
<tr>
<td></td>
<td>$^3$D$_2$</td>
<td>0.0781</td>
<td>0.0790</td>
<td>0.117±0.009</td>
</tr>
<tr>
<td></td>
<td>$^1$D$_2$</td>
<td>0.0430</td>
<td>0.0444</td>
<td>0.072±0.003</td>
</tr>
</tbody>
</table>

Two-electron transitions between discrete levels

K XI $2p^43p \rightarrow 2p^43s$
$2p^43p \rightarrow 2s2p^6$

Radiative lifetimes

<table>
<thead>
<tr>
<th></th>
<th>$\tau_0$</th>
<th>$\tau_1$</th>
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<tbody>
<tr>
<td>$(^1D)3p\ ^2P_{3/2}$</td>
<td>0.119</td>
<td>0.067</td>
</tr>
<tr>
<td>$(^1D)3p\ ^2P_{1/2}$</td>
<td>0.144</td>
<td>0.057</td>
</tr>
<tr>
<td>$(^1S)3p\ ^2P_{3/2}$</td>
<td>0.327</td>
<td>0.147</td>
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<tr>
<td>$(^1S)3p\ ^2P_{1/2}$</td>
<td>0.209</td>
<td>0.175</td>
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</table>

Single electron ($\tau_0$) and two-electron ($\tau_1$) transitions are included
Calculated physical quantities and their approximate accuracy for highly charged ions

- Energy levels: \( \sim 0.4 - 0.2 \% \)
- Wavelengths: \( \sim 0.2 - 0.1\% \)
- Oscillator strengths, transition probabilities: \( \sim 10 - 1 \% \)
- Radiative lifetimes: \( \sim 3 - 1 \% \)
Calculation of spectral lines of tungsten ions

Applying a new Quasi-Relativistic Approach
- for highly charged ions
- for neutral/nearly neutral atoms

Data produced for
- energy level spectra
- radiative transitions, oscillator strengths
- level lifetimes

Mean square deviations $\sigma$ and relative mean-square deviations $S$

<table>
<thead>
<tr>
<th>Method</th>
<th>$W^{29+}$</th>
<th>$W^{30+}$</th>
<th>$W^{31+}$</th>
<th>$W^{32+}$</th>
<th>$W^{33+}$</th>
<th>$W^{34+}$</th>
<th>$W^{35+}$</th>
<th>$W^{29+}$</th>
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</thead>
<tbody>
<tr>
<td>BP</td>
<td>$\sigma$</td>
<td>32100</td>
<td>30000</td>
<td>25200</td>
<td>29100</td>
<td>34965</td>
<td>46400</td>
<td>59200</td>
</tr>
<tr>
<td></td>
<td>$S$</td>
<td>1.89%</td>
<td>1.71%</td>
<td>1.55%</td>
<td>1.48%</td>
<td>1.58%</td>
<td>1.94%</td>
<td>2.50%</td>
</tr>
<tr>
<td>QR</td>
<td>$\sigma$</td>
<td>13700</td>
<td>9800</td>
<td>9800</td>
<td>9000</td>
<td>8700</td>
<td>8900</td>
<td>9000</td>
</tr>
<tr>
<td></td>
<td>$S$</td>
<td>0.73%</td>
<td>0.69%</td>
<td>0.61%</td>
<td>0.51%</td>
<td>0.46%</td>
<td>0.44%</td>
<td>0.47%</td>
</tr>
</tbody>
</table>

Energy level spectra
E1 emission transition probabilities
$E1$ emission transition probabilities

\begin{itemize}
  \item \textbf{W}^{36+}
\end{itemize}
Global characteristics

The essence of the global characteristics of spectra is the replacement of the transitions between separate levels by the transition between configurations.

Usually the spectrum emitted has the shape of broad maximum. Such maximum can be described using the global characteristics of spectra:

- Average energy $E$,
- The variance of transitions between two configurations $\sigma$. 
Global characteristics

They can be expressed in terms of the initial or centered moments

\[ \alpha_k = \sum_i (E_i)^k p_i, \quad \mu_k = \sum_i (E_i - \bar{E})^k p_i, \quad p_i = I_i / I. \]

Here \( I \) is the total intensity of the spectrum, \( I_i \) is the intensity of \( i \)-th line.

The global characteristics depend on the excitation conditions and population of levels.
Global characteristics

Average energy

\[
\bar{E}(K) = \frac{\sum <K\gamma|H|K\gamma>}{g(K)}
\]

\[
k \geq 2 \quad \mu_k(K) = \frac{\sum [<K\gamma|H|K\gamma>-\bar{E} \gamma]^k}{g(K)}
\]

\[
\mu_k(K, K') = \frac{\sum [<K\gamma|H|K\gamma>-K\gamma' H|K\gamma'/>]^k <K\gamma|D|K'\gamma'>^2}{S(K, K')}
\]

\[
k = 2 \quad \text{- dispersion,}
\]
\[
k = 3 \quad \text{- asymmetry coefficient,}
\]
\[
k = 4 \quad \text{- coefficient of the excess}
\]
Thank you