



Gonçalo Garcês S.R. Baptista

Bsc. in Physics Engineering

President: Prof André Wemans

Rapporteur: Prof. Jorge Sampaio Advisor: Prof. Mauro Guerra

Co-Advisor: Prof. Jorge Machado

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X-ray Resonant Raman Scattering

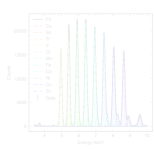
Spectra simulation from first principles for Copper below the ionization threshold using high-performance computing

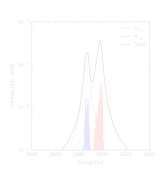


- 1 Theoretical Introduction Characteristic x-rays The Hamiltonian State-of-the-art
- Atomic structure calculations
 The system at study
 Level calculations
 Transition computations
 Fundamental atomic
 parameters
- 3 Spectra simulation
 Line shape
 Photoexcitation
 Photoionization
 The synthetic spectrum
- 4 Spectral Analysis
 The theoretical results
 Comparison with
 experimental data
- A new parallelization code The MPI approach Speedup comparison

X-ray applications





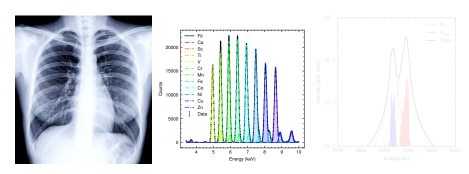


- (a) Imaging purposes

Figure: Application examples of x-ray radiation

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X-ray applications



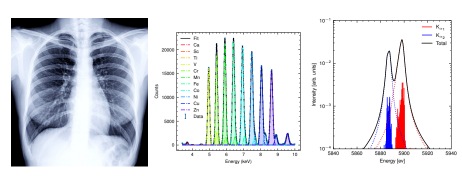
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- (b) Sample quantification (c) Fundamental parameters

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Theoretical Introduction

X-ray applications



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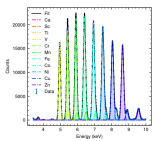
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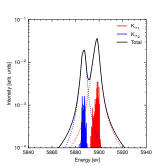
Goncalo Baptista NOVA School of Science and Technology Theoretical Introduction

Characteristic x-rays

X-ray applications







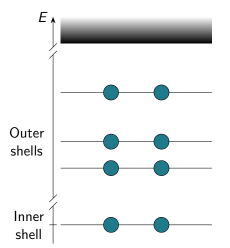
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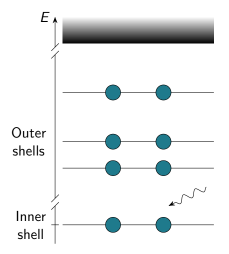
Vacancy generation and relaxation processes



- Bound state system

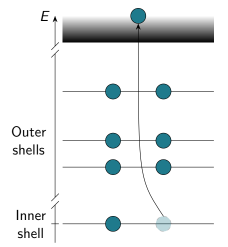
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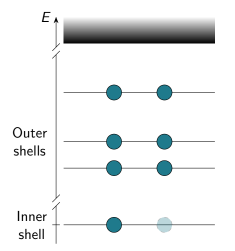
- Bound state system
- Energy transfer
- Ionization
- Vacancy generated
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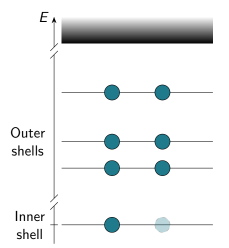
- Ionization





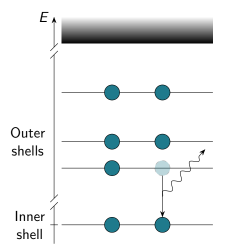
- Vacancy generated





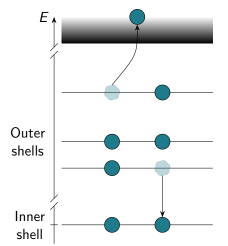
- Vacancy generated
- Atomic Relaxation





- Round state system
- Energy transfer
- Ionization
- Vacancy generated
- Atomic Relaxation
 - Radiative relaxation (x-ray emission)
 - Auger electron emission





- Atomic Relaxation

 - Auger electron emission

Alternative processes

Shake processes

Post-ionization, the different number of particles in the system leads to a change in the Hamiltonian. This leads to the lack of orthogonality between non-equivalent states and free-wave wavefunctions in the initial and final configurations.

$$\langle 1s_i|2s_f\rangle \neq 0$$

$$\langle 1 \mathsf{s_i} \big| e_\mathsf{free}^- \rangle \neq 0$$

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Shake-up

Excitation of extra electron(s).

Theoretical Introduction

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$$\langle 1 \mathsf{s_i} | e_{\mathsf{free}}^- \rangle \neq 0$$

Shake-off

Ionization of extra electron(s).

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Alternative processes

Shake Processes

The probability of these processes occurring is extremely dependent on the incident energy. Since for this thesis, the energy ranges studied were below the ionization threshold, no shake processes were accounted for.

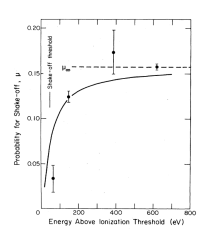


Figure: Thomas model for shake probability. [1]

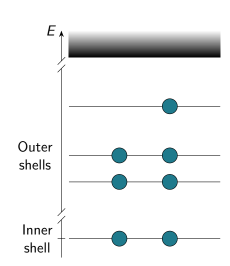
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Other processes

Photoexcitation

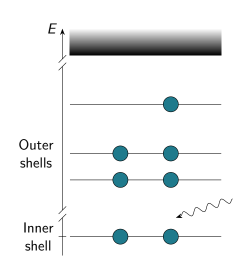
The most relevant process, for the scope of this work, is that of resonant photoexcitation. In it, instead of a vacancy generated due to sending one electron to the continuum, it has, in turn, remained bound but in a higher energy level.



Other processes

Photoexcitation

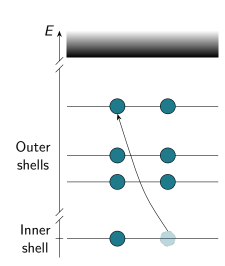
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Theoretical Introduction

The Hamiltonian

Schrödinger's Hamiltonian

In its most basic form, for a "classical" atom (nucleus + electrons), and when relativistic effects are not taken into account, the considered Hamiltonian follows the one use in Schrödinger's equation:

$$\hat{H} = \sum_{i}^{N} \frac{1}{2} \nabla_{i}^{2} - \frac{Z}{r_{i}} + \sum_{j>i} \frac{1}{r_{ij}}$$

- The kinetic energy of the electron.
- The potential energy of the electron-nucleus attraction.
- The potential energy of the electron-electron repulsion

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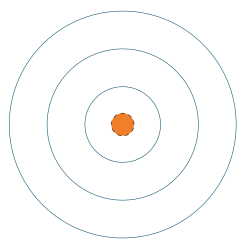
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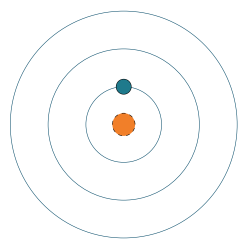
Solving the non-relativistic many-body problem

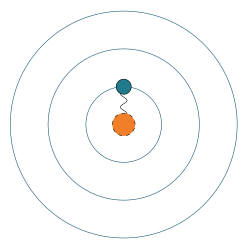
Due to the complexity introduced by the many bodies in the system, and their interactions, a numerical method needs to be employed as to obtain eigenfunctions for this Hamiltonian.

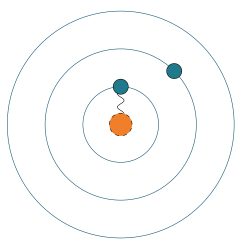


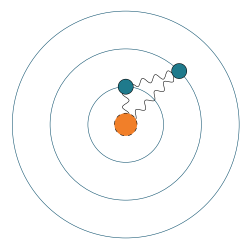
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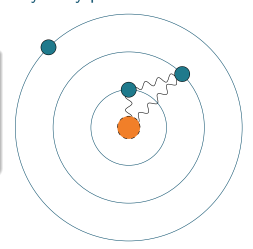
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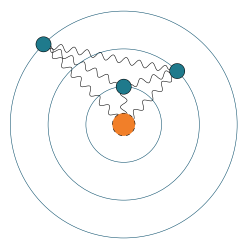












Each of the electrons' wavefunctions, u, are composed as a product of a spatial part, ψ , and one related to the electron's spin χ .

$$u = \psi \chi$$

The system's wavefunction should then be written as a Slater determinant as to account for anti-symetry and the fermionic nature of the electrons.

$$\frac{1}{\sqrt{N!}} \begin{vmatrix} u_1(x_1) & u_2(x_1) & \dots & u_N(x_1) \\ u_1(x_2) & u_2(x_2) & \dots & u_N(x_2) \\ \vdots & \vdots & \ddots & \vdots \\ u_1(x_N) & u_2(x_N) & \dots & u_N(x_N) \end{vmatrix}$$

The Hamiltonian

Solving the non-relativistic many-body problem

Through a self-consistent field approach, the method solves, for each cycle, a set of integro-differential equations as a way to compute new wavefunctions and the new energy for the system.

$$\begin{array}{c|c}
E_{\text{opt}} & & N = \infty \\
E_3 & & N = 4 \\
E_2 & & N = 3
\end{array}$$

$$E_1$$
 — $N = 2$

$$E_0$$
 — $N=1$

The Hamiltonian

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Solving the non-relativistic many-body problem

This process is then repeated up until the energy difference in-between two steps is under a pre-defined benchmark value, as to assure the computation has converged.

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Limitations of the non-relativistic approach

While Schrödinger's equation may be quite accurate for low energy systems (e.g. Hydrogen), where the speed of the surrounding electrons is not comparable to that of light, such is not the case for more complex and heavier systems.

Speed of 1s electrons in ground state configurations (% of c)

Hydrogen: 0.516%

Copper: 14.751%

Uranium: 49.211%

$$E_k = mc^2 \left(\frac{1}{\sqrt{1 - \frac{v^2}{c^2}}} - 1 \right)$$

The Hamiltonian

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Relativistic approaches

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The Hamiltonian

Relativistic approaches

- Incorporates the relativistic

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Dirac equation

- Includes the same advantages as the previous.
- Accounts for spin interactions.
- Yields two-component solutions, for positive and negative energies.
- The "Dirac sea" originated the talks about the possible existence of anti-particles.

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Theoretical Introduction

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Field retardation effects:

Necessary to account for Breit's corrections:

$$H_B = \sum_{i>j} \frac{\mathrm{e}^2}{r_{ij}} - \mathrm{e}^2 \bigg(\frac{\alpha_i \alpha_j}{r_{ij}} + \frac{(\alpha_i \nabla_i)(\alpha_j \nabla_j) r_{ij}}{2} \bigg),$$

Theoretical Introduction

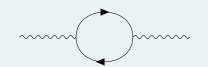
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$$H = H_D + H_B$$

Field quantization effects (QED):

Necessary to account for effects from Quantum Electrodynamics, such as Self-energy and Vacuum Polarization.





Theoretical Introduction

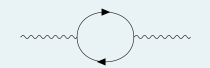
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Theoretical Introduction

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The state-of-the-art

mcdfgme (Multi Configuration Dirac-Fock General Matrix Elements)

This code is a novel computational implementation, based on the Hatree-Fock method, capable of calculating a plethora of atomic parameters, while incorporating all previously mentioned necessary considerations.

It has proven time and time again to have excellent accuracy and precision when performing calculations for the most varied systems.

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Overview

- Theoretical Introduction Characteristic x-rays
- Atomic structure calculations The system at study Level calculations Transition computations Fundamental atomic parameters
- Spectra simulation
- 4 Spectral Analysis
- **6** A new parallelization code

The considered excitations

Atomic structure calculations

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As previously stated, the focal point of this work is that of computing radiative relaxation spectra for excited Copper.

Ground State:
$$1s^2 \ 2s^2 \ 2p^6 \ 3s^2 \ 3p^6 \ 3d^{10} \ 4s^1$$

Excitations of any one of the consituent electrons to orbitals:

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The calculated configurations

In order to account to all possible decay possibilities, it was necessary to perform calculations for two different sets of configurations.

1-hole configurations

Related to the initial and final states of radiative transitions. Obtained by running a hole through all orbitals in the base-configurations.

2-holes configurations

Related to the final states of Auger transitions. Obtained by running a combination of two holes through

all orbitals in the base-configurations.

The level manifold

It is now necessary to compute all possible levels the studied atomic system can be in. As it will be shown, this is no simple task...

The level manifold

Besides the original configuration, three other sets of quantum numbers are necessary for defining a level:

- The hole orbital labels. $(n l_i)$
- The total angular momentum number.
- The eigenvalue/Lagrange multiplier.

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Here, we know two base things:

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- One of the electrons was excited to 4p

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This leads to the possibility of the electron having come from:

- 1s
- 2s
- 2p
- 3*s*
- 3p
- 3d
- 45

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These are the labels $(n \mid l_i)$. 7 in total.

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The level manifold- an example: $2p \rightarrow 4p$ excited Copper

For this configuration, three subshells are open: 2p, 4s, and 4p.

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For this configuration, three subshells are open: 2*p*, 4*s*, and 4*p*.

Atomic structure calculations

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The different angular momentum couplings lead to various values for system's total angular momentum.

Possibilities for J values follow:

2p		4s		4p		Total/J
M_I	M_s	M_I	M_s	M_I	M_s	Total/ J $M_I + M_s$
1	-1/2	0	-1/2	1	-1/2 $-1/2$	1/2
1	1/2				-1/2	3/2
1	$1/_{2}$	0	$1/_{2}$	1	-1/2	5/2
1	1/2	0	1/2	1	$1/_{2}$	7/2

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Possibilities for J values follow:

2p		4s <i>M_I M_s</i>		4p		Total/J
M_I	M_s	M_I	M_s	M_I	M_s	$M_I + M_s$
1	_1/2	0	_1/2	1	-1/2 $-1/2$	1/2
1	1/2	0	-1/2	1	-1/2	3/2
1	$1/_{2}$	0	$1/_{2}$	1	-1/2	5/2
_ 1	1/2	0	$1/_{2}$	1	1/2	7/2

Level calculations

The level manifold- an example:
$$2p \rightarrow 4p$$
, $J = 5/2$

Even for this **very** specific example, the branching-out continues:

Level calculations

The level manifold- an example: $2p \rightarrow 4p$, J = 5/2

There are many possibilities for achieving this certain combination of a given configuration and J value. Each, is represented by the respective ϵ .

The level manifold- an example: $2p \rightarrow 4p$, J = 5/2

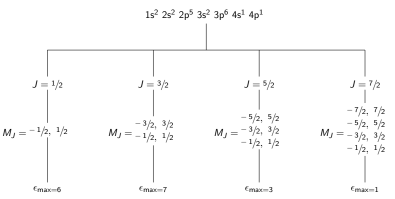
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J = 5/2					
2p		4s		4p	
M_I	M_s	M_I	M_s	M_I	M_s
1	-1/2	0	1/2	1	1/2
1	$1/_{2}$	0	-1/2	1	$1/_{2}$
1	1/2	0	1/2	1	-1/2

The level manifold

This set of possibilities and rearrangements form what we call the level manifold. In addition, a 2J + 1 level degeneracy is accounted for.



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Level calculations with mcdfgme

A calculation was performed for each existent level.

In each of them, a level was treated as a linear combination of state wavefunctions (associated with the eigenvalues) with mixing coefficients:

$$|\Psi\rangle = a_1 |\psi_1\rangle + a_2 |\psi_2\rangle + \dots$$

Self-consistent field

- Coulomb interactions
- Breit considerations
- Vacuum Polarization

Perturbation theory

Self-energy

Level calculations

Evaluating the calculation

What to look out for:

- Orthogonality conservation
- Energy divergences
- Effective nuclear charge

Evaluating the calculation

What to look out for:

- Orthogonality conservation
- Energy divergences
- Effective nuclear charge

Due to the method's numerical nature, orthogonality problems may surge.

We look at overlap values for similar orbitals (same l_j) and set a maximum threshold:

$$|\langle n | I_j | m | I_j \rangle| \le 10^{-6}$$

Evaluating the calculation

What to look out for:

- Orthogonality conservation
- Energy divergences
- Effective nuclear charge

A component of the energy is computed through two different methods. For good convergence, the difference in their values should not be above 1 eV.

$$|E_1-E_2|\leq 1 \text{ eV}$$

Evaluating the calculation

What to look out for:

- Orthogonality conservation
- Energy divergences
- Effective nuclear charge

The presence of inner electrons leads to a shielding of the positive nuclear charge. As so, outer electrons will be subject to an attenuated positive charge. This effect has been previously studied, and the obtained values should be benchmarked.

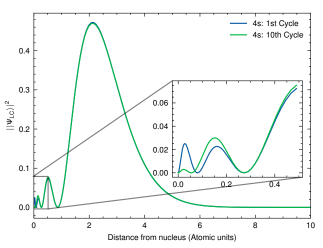
Level calculations

Methods for solving the convergence problems

- Changing the number of self-consistent cycles.
- Altering cycle parameters, such as accuracy.
- Choosing the initial trial wavefunctions for select orbitals (e.g. Hydrogenoids or obtained through the Thomas Fermi potential).
- Changing the method for solving the Dirac equation.
- Enforcing the node number for the wavefunctions

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Changes in wavefunctions after the variational process



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Total number of calculated levels

For the 19 performed calculations,

- 1601 1-hole levels
 - 166 manually converged
- 20550 2-holes levels
 - 1678 manually converged

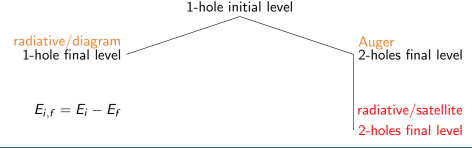
Leaving a total of:

- 22151 levels calculated
- 1844 levels manually converged

Atomic transitions

When a system is not found in its least energetic state various processes will occur until the most stable one is reached.

In this way, for every level, calculations were performed for decays to all possible lower energy states.



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Radiative transitions

Given the set of all 1-hole levels, with n elements, the total number of these transitions (without accounting for level degeneracy), N, is simply given by the amount of combinations of two elements between the set:

$$N = \frac{n!}{2 \cdot (n-2)!}$$

For these transitions, full orbital relaxation was allowed.

Atomic structure calculations

The *mcdfgme* calculation yields the rates for each Electric and/or Magnetic component/pole for both Coulomb (length) and Lorentz (velocity) gauges.

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Auger transitions

The total number of Auger transitions can not be calculated a priori. Since the 1-hole and 2-holes level sets are "independent", the level structure needs to be calculated in order to fully resolve the number of transitions. For computing these transitions, the free-electron wavefunction has to be computed for the initial state potential, and orthogonality is enforced between the free-wave and bound orbitals.

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All calculated transitions

- 70885 diagram radiative



Atomic structure calculations

All calculated transitions

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All calculated transitions

Atomic structure calculations

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- 70885 diagram radiative
- 452988 Auger

All calculated transitions

- 70885 diagram radiative
- 452988 Auger
- 6684258 satellite radiative

Transition computations

All calculated transitions

In total, for this work.

- 70885 diagram radiative
- 452988 Auger
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transitions were calculated, totaling to 7208131 computations performed.

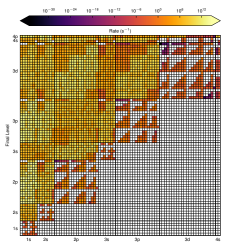
Rate and Energy matrices

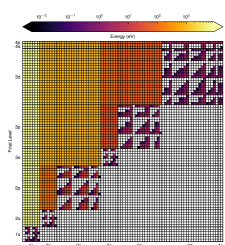
To evaluate and benchmark a calculation, a visualization tool was used, where the calculated energy and rate values were displayed in a grid-like view.

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Transition computations

Rate and Energy matrices

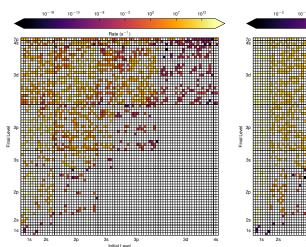


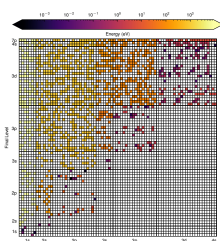


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Rate and Energy matrices





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Initial Level

X-ray Resonant Raman Scattering

Total radiative and non-radiative rates

After computing all possible transitions, one can now start to evaluate certain atomic parameters.

Total radiative and non-radiative rates

To start, the total radiative and non-radiative rates for each level should be computed. When compared, they can provide information on the probability of said level decaying either way.

$$R_i^R = (2J_i + 1) \sum_f R_{i,f}^R$$

$$R_i^{NR} = (2J_i + 1) \sum_{\varepsilon} R_{i,f}^{NR}$$

Fundamental atomic parameters

Fluorescence Yield

The quantity is an indicator of the probability of a level decaying either via the emission of a photon as opposed to an Auger electron.

It is given by the ratio between the level's radiative and total rates:

$$\omega_i = \frac{R_i^R}{R_i^R + R_i^{NR}}$$

Fluorescence Yield

For ionized Copper, the *K* shell F.Y. was computed to be 45.34%. Every calculated excited state presented a value inferior to this.

Minimum value:

6*s*-excited Copper 18.09%

Maximum value:

4*s*-excited Copper 44.88%

This decrease can be explained by the presence of an extra electron increasing the number of Auger channels.

Fluorescence Yield

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Other subshells

For other subshells, save for $M_{4,5}$, the F.Y. slightly oscillated around the obtained value for ionized Copper, but mostly preserving the same order of magnitude.

$M_{4.5}$ subshells

Curiously both for ionized Copper and for all excitations to ns orbitals, as well as 4p and 5p, only radiative decays occur, while for the rest, Auger transitions reign.

Quantum mechanics tells us bound systems are arranged in states with quantized energy, hence why a discrete amount of levels was calculated, a not a continuum.

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It does, however, also alert us to natural uncertainties, as described by the well-known Heisenberg uncertainty principle.

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While the momentum-position relation is certainly the most famous one, its energy-time counterpart will have implications in the next steps:

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While the momentum-position relation is certainly the most famous one, its energy-time counterpart will have implications in the next steps:

$$\Delta E \Delta t \geq \hbar$$

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Fundamental atomic parameters

Level widths

$$\Delta E \Delta t > \hbar$$

From this, we can naturally deduce that, a level which decays to other lower energy ones $(R \to \frac{1}{\Lambda t})$, should, in principle, also have an energy width.

As such, we define the level width as:

$$\Gamma_i = \hbar \cdot \left(R_i^R + R_i^{NR} \right)$$

$$\Delta E \Delta t > \hbar$$

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As such, we define the level width as:

$$\Gamma_i = \hbar \cdot \left(R_i^R + R_i^{NR} \right)$$

And the width of a transition as the sum of the widths for the initial and final level:

$$\Gamma_{i,f} = \Gamma_i + \Gamma_f$$

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Transition intensities

Now that many atomic parameters are calculated, it is possible to calculate the spectral intensity of a given transition.

$$I_{i,f} = N_i \frac{2J_i + 1}{g_{\text{sub}}} \frac{R_{i,f}}{R_i^R + R_i^{NR}}$$

Put into simple terms, the intensity is nothing more than the product of the statistical weight of the level multiplicity, of the individual rate compared to the total one, and a scaling factor accounting for population generation.

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Overview

- Theoretical Introduction
 Characteristic x-rays
 The Hamiltonian
 State-of-the-art
- Atomic structure calculations
 The system at study
 Level calculations
 Transition computations
 Fundamental atomic
 parameters
- Spectra simulation Line shape Photoexcitation Photoionization The synthetic spectrum
- 4 Spectral Analysis
 The theoretical results
 Comparison with
 experimental data
- A new parallelization code The MPI approach Speedup comparison

Line shape

The theoretical spectral shape

Based on all previously computed atomic parameters, it is now possible to start simulating spectral lines.

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Lorentzian profile

Describes the theoretical line shape for the emission:

$$\frac{I_{i,f}}{2\pi} \frac{\Gamma_{i,f}}{(E - E_{i,f})^2 + (\Gamma_{i,f}/2)^2}$$

Does not account for thermal distributions and stochastic events.

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Gaussian profile

This profile will be used to account for any broadening effects that may impact the measured spectra.

$$\frac{1}{\sigma\sqrt{2\pi}}\exp\left(-\frac{E^2}{2\sigma^2}\right)$$

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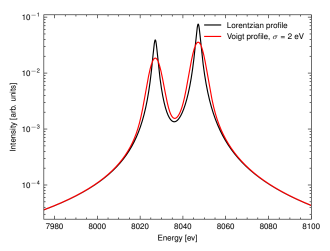
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$$\frac{1}{\sigma\sqrt{2\pi}}\exp\left(-\frac{E^2}{2\sigma^2}\right)$$

The convolution of both gives rise to the Voigt profile, incorporating all the desired effects, at the cost of computational power.

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X-ray Resonant Raman Scattering



The incident beam

For the purpose of this work, the desired source of radiation for such study would be a synchrotron line.

- Due to the usage of wigglers and undulators, the beam can be close to monochromatic.
- The energy of the radiation used is tunable, allowing for the survey from regions below to post-ionization threshold.

As such, a beam profile of 0.5 eV was considered.

What about N_i ? The photoexcitation case

The computation of this parameter is somewhat tricky.

- No analytical expression for cross-sections could be found.
- Should be calculated ab-initio (no data from outside).
- Must simulate the resonance effect.
- Needs to be valid (according to the laws of physics).
- The beam profile should be considered.



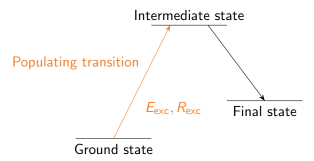


What about N_i ? The photoexcitation case

The proposed solution:

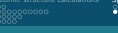
To solve this problem, an ad-hoc expression was developed.

The rate and energy of the transition from ground state to the excited state was considered by looking at the inverse transition.



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What about N_i ? The photoexcitation case

A Lorentzian profile was then created using both the energy value, and the associated rate. The overlap between this profile and the beam's was computed and multiplied by the rate:

$$N_i = R_{\mathsf{exc}} \int_{-\infty}^{\infty} G(E - E_{\mathsf{beam}}, 0.5 \; \mathsf{eV}) \cdot L(E - E_{\mathsf{exc}}, \Gamma_{\mathsf{exc}}) \; \mathrm{d}E$$

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Conclusion

Thank you for your attention!

