Abstract: The online atomic database of NORAD-Atomic-Data, where NORAD stands for Nahar OSU Radiative, is part of the data sources of the two international collaborations of the Opacity Project (OP) and the Iron Project (IP). It contains large sets of parameters for the dominant atomic processes in astrophysical plasmas, such as, (i) photo-excitation, (ii) photoionization, (iii) electron–ion recombination, (iv) electron–impact excitations. The atomic parameters correspond to tables of energy levels, level-specific total photoionization cross-sections, partial photoionization cross-sections of all bound states for leaving the residual ion in the ground state, partial cross-sections of the ground state for leaving the ion in various excited states, total level-specific electron–ion recombination rate coefficients that include both the radiative and dielectronic recombination, total recombination rate coefficients summed from contributions of an infinite number of recombined states, total photo-recombination cross-sections and rates with respect to photoelectron energy, transition probabilities, lifetimes, collision strengths. The database was created after the first two atomic databases, TOPbase under the OP and TIPbase under the IP. Hence the contents of NORAD-Atomic-Data are either new or from repeated calculations using a much larger wave function expansion making the data more complete. The results have been obtained from the R-matrix method using the close-coupling approximation developed under the OP and IP, and from atomic structure calculations using the program SUPERSTRUCTURE. They have been compared with available published results which have been obtained theoretically and experimentally, and are expected to be of high accuracy in general. All computations were carried out using the computational facilities at the Ohio Supercomputer Center (OSC) starting in 1990. At present it contains atomic data for 154 atomic species, 98 of which are lighter atomic species with nuclear charge \( Z \leq 28 \) and 56 are heavier ones with \( Z > 28 \). New data are added with publications.

Keywords: atomic data; transition probabilities; photoionization cross sections; electron–ion recombination cross sections and rate coefficients; lifetimes; electron–impact excitation

1. Introduction: NORAD-ATOMIC-DATA

The online atomic database NORAD-Atomic-Data [1] was established at the Ohio State University in 2007 at the suggestion of astronomers and initiative of Dr. C. Allende Prieto. The database is accessible from links provided at various database pages, such as International Atomic Energy Agency [2], CfA-Harvard University [3]. Current data at NORAD-Atomic-Data are for 154 atoms and ions, presented on the website [1] in two tables, Table 1 containing 98 ions with \( 1 \leq Z \leq 28 \) and Table 2 55 ions with \( 28 < Z \leq 92 \) from published papers. All data files at NORAD-Atomic-Data are in standard ASCII format. They can be downloaded for direct use in models and for diagnostics of astrophysical and laboratory plasmas.

A large part of the data in NORAD-Atomic-Data was obtained from research under two international collaborations, The Opacity Project (OP) [4] (UK, USA, Germany, France, Venezuela, ...
Canada) and Iron Project (IP) [5] (UK, USA, Germany, France, Belgium, Venezuela). The OP [4] ran during 1983–2007 and restarted in 2011, and the IP [5] started in 1993 and is continuing. The aims of the OP and IP are to (i) study in detail the atomic radiative and collisional processes of elements that are abundant in astrophysical plasmas, (ii) compute large volume high precision atomic data with consideration of a large number of energy levels, typically \( n \leq 10 \), and (iii) apply the computed data to solve long-standing astrophysical problems, such as stellar plasma opacity. NORAD-Atomic-Data is the follow-up of the two atomic databases. TOPbase [6,7], which provides energy levels, oscillator strengths and photoionization cross sections in LS coupling, was obtained for the first time for many atoms and ions under the OP. The second database is the TIPbase [8] which provides collision strengths and atomic data for the radiative processes of a limited number of atomic systems obtained with fine structure.

NORAD contains a vast number of parameters for the four dominant atomic processes, which are described below, involving the interaction of a photon or electron and an ion that contribute the most to the spectra of astrophysical and laboratory plasmas. These parameters have been obtained mainly from the R-matrix method using R-matrix package of codes developed under the OP and IP [9–11] and their extensions for theoretical spectroscopy in LS coupling (e.g., [12]), for fine structure splitting of transitions (e.g., [13]), for fine structure spectroscopy (e.g., [14,15]), from the unified method for electron–ion recombination calculations (e.g., [16–18]), and from atomic structure calculations using the code SUPERSTRUCTURE (SS) [19,20]. The large-scale computations are carried out using high-performance computational facilities at the Ohio Supercomputer Center (OSC) where the codes were installed by the OSU group of the OP and the IP team in 1990. The atomic processes and the relevant atomic quantities available at NORAD-Atomic-Data are described below:

1.1. Photoexcitation and De-Excitation
These processes of an ion \( X^{+Z} \) of charge \( Z \) (zero for a neutral) are:

\[
X^{+Z} + h\nu \rightleftharpoons X^{+Z*}
\]  

(1)

The relevant parameters are
- Line strength (\( S \)),
- Oscillator strength (\( f \)),
- Radiative decay rate (\( A \)-value)
- Lifetimes

The database contains data for transitions of the following types: allowed electric dipole (E1) of the same (LS-allowed) and different (intercombination) spin multiplicities, electric quadrupole (E2), electric octupole (E3), magnetic dipole (M1), and magnetic quadrupole (M2). The data for E1 transitions are obtained mainly from the R-matrix method and its extensions while those for the forbidden transitions have been obtained from Breit–Pauli atomic structure calculations using code SS. Individual transitions form spectral lines, however, data for a large number of transitions are needed to determine the opacity in plasmas, modeling synthetic spectra.

1.2. Photoionization
Photoionization (PI) proceeds directly as

\[
X^{+Z} + h\nu \rightarrow X^{+Z+1} + e
\]  

(2)

which provides only the background cross section, and indirectly via formation of an intermediate doubly-excited autoionizing state as

\[
h\nu + X^{+Z} \rightleftharpoons (X^{+Z})^{**} \rightleftharpoons X^{+Z+1} + e.
\]  

(3)
This indirect process introduces a resonance as well as an impact on the background cross-section. The relevant parameters are:

- Total photoionization cross-section ($\sigma_{PI}$) of each bound state of the ion, from ground to various excited states, typically with $n \leq 10$. Total $\sigma_{PI}$ corresponds to the summed contribution of all ionization channels leaving the residual ion in the ground and various excited states.
- Partial photoionization cross-section ($\sigma_{PI}$) for all bound states of the ion for leaving the residual ion in the ground state only.
- Partial photoionization cross-sections ($\sigma_{PI}$) corresponding to leaving the residual ion in the ground and various excited states. NORAD-Atomic-Data provides these cross-sections only for the ground level of the ion. Photoionization resonances are often dissolved by plasma density and temperature, resulting in an enhanced continuum background. However, the strong and isolated resonances can be seen in the absorption spectra. Large quantities photoionization cross-section for all possible bound states with a wide range of photon energies are needed to determine the opacity in astrophysical plasmas.

1.3. Electron–Ion Recombination

Electron–ion recombination is the inverse process of photoionization. It can proceed directly, known as radiative recombination (RR), as

$$X^{+Z+1} + e \rightarrow X^{+Z} + h\nu,$$

which provides only the background recombination rate, and indirectly through an autoionizing state, known as dielectronic recombination (DR), as

$$e + X^{+Z} \Leftrightarrow (X^{+Z-1})^{**} \Leftrightarrow \begin{cases} e + X^{+Z} \quad \text{AI} \\ X^{+Z-1} + h\nu \quad \text{DR} \end{cases}$$

AI is autoionization, the other decay channel of an autoionizing state leading to ionization instead of radiative stabilization.

Relevant parameters for the process obtained from the unified method of Nahar and Pradhan [16,17] are:

- Level-specific total recombination (includes both RR and DR) rate coefficients ($\alpha_{RC}(i)$) of all bound levels with $n \leq 10$.
- Total recombination rate coefficient ($\alpha_{RC}(T)$), summed contributions of all levels with $n \leq 10$ and of levels with $10 < n \leq \infty$ as function of electron temperature $T$.
- Total recombination cross sections ($\sigma_{RC}(E)$) and total recombination rate coefficient ($\alpha_{RC}(E)$) with respect to photoelectron energy $E$. Recombination resonances can be seen in emission spectra as dielectronic recombination lines. Total $\alpha_{RC}(T)$ is needed for the determination of ionization fractions in photoionized or collisional plasmas.

1.4. Electron–Impact Excitation (EIE)

EIE describes exciting an electron of the target ion by impact of a projectile electron. The state then drops down by giving out a photon that carries the information of the excitation of the target.

$$e + X^{+Z} \rightarrow e' + X^{+Z*} \rightarrow X^{+Z} + h\nu + e'.$$

The excitation can be an autoionizing state which includes a decay channel for autoionization. Relevant parameters are:

- Collision strength ($\Omega$),
- Effective collision strengths ($\gamma$),
Collision rate coefficients ($q_{ij}$).

Emission lines from levels excited by EIE provides diagnostics of thin plasmas. Although largely similar, NORAD-Atomic-Data has some differences from TOPbase and TIPbase as explained below.

1.5. The Opacity Project and atomic database TOPbase

The Opacity Project [4] was initiated in 1983 at the plea of astronomers due to the need for accurate atomic data. The comparison of observed and predicted astrophysical spectra suggested a few times greater abundance of metals (elements heavier than Li) compared to model predictions. Under the OP leadership of Michael J. Seaton, the original R-matrix codes (e.g., for collisions [21]) went through massive changes with huge efforts, the introduction of detailed photoionization with resonances. Systematic studies of radiative processes of photoexcitations and photoionization with resonances were carried out for all astrophysically abundant atoms and ions from hydrogen to iron for the first time by a team of 25 investigators in Europe, US, and South America. Extensive volumes of atomic data they computed under the OP by were made available through the database TOPbase in 1993 [6,7]. The results were obtained in LS coupling, so the fine structure splitting of energy levels was not accounted for. However, the effort solved and continues to solve many astrophysical problems and has revealed new features in photoionization.

It may be added that monochromatic opacities of elements from H to Fe obtained using the atomic data of TOPbase are available at the OP Server [22] at OSC.

1.6. The Iron Project and atomic database TIPbase

Following the crucial role played by the OP for radiative atomic processes, the need for high-accuracy collisional atomic data was realized. This initiated the Iron Project in 1991 to study electron impact excitation (EIE) with high accuracy for iron and iron-peak elements that had very limited data due to difficulty in computation arising from their complex atomic structure. In addition to EIE, the IP included computing radiative atomic data with inclusion of the next level of accuracy with fine structure and other relativistic effects. The R-matrix codes were further extended to include relativistic effects in the Breit–Pauli approximation, which was named as the Breit–Pauli R-matrix or Breit–Pauli R-matrix (BPRM) method [5,11]. Extensive volumes of atomic data have been obtained, and continue to be computed, under the IP. The resulting atomic data formed the database TIPbase [8]. However, this database contains only a part of the work carried out under the IP.

1.7. NORAD-ATOMIC-DATA and Atomic Astrophysics and Spectroscopy

The extensive effort at the Oho State University (OSU) resulted in the online database, NORAD-Atomic-Data [1]. It contains data mainly for the radiative processes and limited for the collisional process, obtained beyond TOPbase. and are not available in TOPbase or TIPbase. It has some differences in structure and contents from those of TOPbase and TIPbase. Atomic data in NORAD-Atomic-Data have been obtained by Nahar et al. only, and it is maintained by Nahar, the present author. The main differences are as follows:

- Significant part of the data corresponds to new and/or improved results over those in TOPbase,
- The data are of higher accuracy than those in TOPbase,
- Includes fine structure oscillator strengths of many ions,
- Contains forbidden transitions, lifetimes,
- Includes data of the additional atomic process of electron–ion recombination,
- Contains larger sets (typically up to $n = 10$ and $l \leq 9$) of energy levels, photoionization cross-sections, recombination cross sections and rate coefficients needed for complete modeling of astrophysical objects,
- Contains X-ray transition data for heavier elements (beyond Ni) which are of great interest for various astronomical, biomedical, fusion plasma applications.
The textbook “Atomic Astrophysics and Spectroscopy” [23] is a culmination of findings in atomic processes revealed in the study under the OP and IP. It describes details of atomic physics and their relevance to astronomical applications, interpretation of astrophysical spectra with atomic processes, and forms the bridge between atomic physics and astronomy.

2. Theoretical Approximations

The theoretical background used for obtaining the atomic parameters gives a measure of general consistency and accuracy of the results. Hence a brief outline of the theory and description of definitions of quantities are given here to inform data users. While most of the results in NORAD-Atomic-Data have been obtained from ab initio R-matrix method using the close-coupling (CC) approximation (e.g., [9,21,23]), transition probabilities have also been obtained from atomic structure calculations using the SUPERSTRUCTURE (SS) software. SS is also a part of the R-matrix package of code, providing the initial wave function of the core (also termed as residual or target) ion to initiate R-matrix calculations.

The relativistic Hamiltonian in the BPRM method adopted under the IP is given by ([20,23])

\[ H_{BP} = H_{NR} + H_{mass} + H_{Dar} + H_{so} + \sum_{i \neq j}^{N} \left[ g_{ij}(so + so') + g_{ij}(ss') \right] \]

(7)

The non-relativistic Hamiltonian is

\[ H_{NR} = \left\{ \sum_{i=1}^{N} \left\{ -\nabla_i^2 - \frac{2Z}{r_i} + \sum_{j>i}^{N} \frac{2}{r_{ij}} \right\} \right\} \]

(8)

where \( i, j \) are indices for electrons, \( r \) is the distance, and \( Z \) is the nuclear charge of the ion. The one-body relativistic correction terms are \( H_{mass} = -\frac{\hbar^2}{2m} \sum_{i} p_i^4 \), \( H_{Dar} = \frac{\hbar^2}{2m} \sum_{i} \nabla^2 \left( \frac{Ze}{r_i} \right) \), \( H_{so} = \frac{Ze^2 \hbar^2}{2m^2c^2} \mathbf{L} \cdot \mathbf{S} \), where \( \alpha \) is the fine structure constant, \( p_i \) is the momentum of electron \( i \), \( m, e \) are electron mass and charge, \( h, c \) are constants, and \( \mathbf{L}, \mathbf{S} \) are the total orbital and total spin angular momenta of the ion. The two-body Breit interaction is \( H_{B} = \sum_{i>j} \left[ g_{ij}(so + so') + g_{ij}(ss') \right] \) where various \( g \) represent terms of spin–orbit \( so \), spin–other orbit \( so' \), spin–other spin \( ss' \) interaction [23]. BPRM includes relativistic contributions from these terms and part of other two-body interaction terms [23].

Wave functions and energies are obtained solving:

\[ H \Psi = E \Psi \]

(9)

in the CC approximation, where the atom/ion described as an \((N + 1)\)-electrons system, the target or the core ion is an \(N\)-electrons system, which is interacting with \((N + 1)\)th electron. The total wave function is an expansion (e.g., [9,23])

\[ \Psi_{E}(e + \text{ion}) = A \sum_{i}^{N} \chi_{i}(\text{ion})\theta_{i} + \sum_{j} \Phi_{j}(e + \text{ion}), \]

(10)

where \( \chi_{i} \) represents the core ion wave function, which is typically obtained from SS, \( \theta_{i} \) represents the interacting electron wave function, which can be free at energy \( E \geq 0 \) or bound at \( E < 0 \). The sum on \( i \) represents the number of core ion states included in the R-matrix calculations. \( A \) is the anti-symmetrization operation. \( \Phi_{j} \) in the bound-channel term (the second term) are the correlation configurations of the \((e + \text{ion})\) system and number of \( j \) values depends on the choice of configurations that might contribute significantly for the improvement of energies of the ion.

Complex resonance structures in atomic processes are introduced via channel couplings of the continuum (free) and bound wave function. An electron in a Rydberg state \( E_{ex} \nu l > 0 \), where \( E_{ex} \) is an excited core ion energy in Ry and \( \nu l \) are the effective quantum number and orbital angular momentum.
of the interacting electron, introduces a Rydberg resonance at \( E_{xn} = E^{\ast\ast}(X^+ \nu l) = E_x - z^2/\nu^2 \)
where \( z \) is the charge of the core ion. The other type of resonance, Seaton’s resonance due to PEC (photo-excitation-of-core) appears when the incident photon energy \( h\nu \) is equal to \( E_x \) for a dipole-allowed transition in the core ion. Please note \( \nu \) with a photon represents photon frequency and is different from effective quantum number \( \nu \) that appears with angular momentum. The feature of the CC wavefunction which can form resonances naturally is not available in other approximations, e.g., distorted wave Born approximation (DWBA).

Substitution of \( \Psi_E(e + i\text{on}) \) expansion in the Schrodinger equation results in a set of coupled equations, which are solved by the R-matrix method.

In the R-matrix method, the space is divided into two regions, the inner and the outer regions, of a sphere of radius \( r_a \) with the ion at the center as illustrated in Figure 1. \( r_a \) is large enough to include electron-electron interaction potential inside and is almost zero outside the boundary. Hence outside potential at \( r > r_a \) is Coulombic from perturbation of long-range multipole potentials. In the inner region, the radial part \( F_i(r)/r \) of the outer electron wave function (\( \theta \)) is expanded in terms of a basis set, known as the R-matrix basis,

\[
F_i = \sum_k a_k u_k(r_i)
\]

that satisfies

\[
\left[ \frac{d^2}{dr^2} - \frac{l(l+1)}{r^2} + V(r) + \epsilon_{lk} \right] u_{lk} + \sum_n \lambda_{nlk} P_{nl}(r) = 0
\]

and is made continuous with Coulomb functions outside \( r_a \) (see for example [23]). \( V(r) \) is the potential experienced by the interacting electron, \( \epsilon_{lk} \) is the energy eigenvalue, and \( \lambda_{nlk} \) are Lagrange multipliers. \( P_{nl}(r) \) is a bound orbital wave function of the target. Computations using BPRM method involve a number of stages as described in the flowchart of Figure 2.

As shown in Figure 2, R-matrix calculations can have 3 branches to proceed—(1) LS coupling and relativistic Breit–Pauli, (2) large configuration interaction LS coupling, (3) Dirac relativistic. Various stages generate Hamiltonian matrix (\( H \)), dipole matrices (\( D \)), bound wave functions (\( B \)), and continuum wave functions (\( F \)). End results are (1) energy levels, (2) oscillator strengths, (3) photoionization cross sections, (4) recombination rate coefficients, and (5) collision strengths, which are applied to astrophysical models.

For atomic structure calculations using SUPERSTRUCTURE, the wave functions and energies are obtained using Thomas–Fermi–Dirac–Amaldi potential. The potential represents atomic electrons as a
Fermi sea, constrained by the Pauli exclusion principle, filled in cells up to a the highest Fermi level of momentum:

\[ V(r) = \frac{Z_{\text{eff}}(\lambda_{nl}, r)}{r} = -\frac{Z}{r} \phi(x), \]  

where \( \phi(x) = e^{-2x/2} + \lambda_{nl}(1 - e^{-2x/2}) \), \( x = e^{-Zr/2} \), \( \mu = 0.8853 \left( \frac{N}{N-1} \right)^{2/3} Z^{-1/3} \) is a constant. The function \( \phi(x) \) is a solution of the potential equation [19]. The quantities \( \lambda_{nl} \) are the scaling parameters for the orbitals.

With the wavefunctions, various atomic quantities can be calculated as given below.

\[ f_{ij} = \left[ \frac{E_{ji}}{3g_i} \right] S, \quad A_{ji}(\text{sec}^{-1}) = \left[ 0.8032 \times 10^{10} \frac{E_{ji}^3}{3g_j} \right] S \]  

Figure 2. Flowchart of different stages, e.g., STG1, STG2, RECUPD, of computation in the Breit–Pauli R-matrix (BPRM) method. The final product parameters are energy levels, oscillator strengths, photoionization cross sections, electron–ion recombination cross sections, collision strengths which are applied in astrophysical applications.

2.1. Photoexcitations/Deexcitations

The radiative transition matrix, \( T_{BB'} = \langle \Psi_B || D || \Psi_{B'} \rangle \) for dipole allowed transitions with initial and final bound wavefunctions \( \Psi_B \) and \( \Psi_{B'} \), and the dipole operator \( D = \sum n \mathbf{r}_n \) where \( n \) is the number of electrons, provides the line strength

\[ S = \left| \langle \Psi_f || D || \Psi_i \rangle \right|^2 = \left| \left( \Psi_f \sum_{j=1}^{N} r_j \Psi_i \right) \right|^2 \]  

where \( \Psi_i \) and \( \Psi_f \) are initial and final wave functions. Oscillator strengths \( (f) \) and the transition probabilities \( (A \text{-values}) \) for E1 transitions are obtained as
where \( E_{ji} \) is the transition energy and \( g_i \) is the statistical weight factor. The forbidden transitions obtained using SS are of type (i) electric quadrupole (E2) and (ii) magnetic dipole (M1),

\[
i) \ A_{ji}^{E2} = 2.6733 \times 10^5 \frac{E_{ji}^5}{g_j} S^{E2}(i,j) \text{ s}^{-1}, \quad ii) \ A_{ji}^{M1} = 3.5644 \times 10^4 \frac{E_{ji}^3}{g_j} S^{M1}(i,j) \text{ s}^{-1},
\]

(iii) Electric octupole (E3) and (iv) magnetic quadrupole (M2),

\[
iii) \ A_{ji}^{E3} = 1.2050 \times 10^{-3} \frac{E_{ji}^7}{g_j} S^{E3}(i,j) \text{ s}^{-1}, \quad iv) \ A_{ji}^{M2} = 2.3727 \times 10^{-2} s^{-1} \frac{E_{ji}^5}{g_j} S^{M2}(i,j).
\]

### 2.2. Photoionization (PI)

The radiative transition matrix for photoionization, \( T_{BF} = \langle \Psi_f | D | \Psi_i \rangle \) is obtained from bound and continuum wave functions which give the line strength using the expression above. Photoionization cross section is obtained as

\[
\sigma_{PI} = \frac{4\pi}{3c} \frac{1}{g_i} \omega S,
\]

where \( \omega \) is the incident photon energy in Rydberg unit.

### 2.3. Electron–Ion Recombination

Under the unified method, recombination cross section, \( \sigma_{RC} \), for all bound levels up to \( n \leq n_o \) (typically \( n_o = 10 \)) is obtained from the principle of detailed balance (Milne relation) as

\[
\sigma_{RC} = \sigma_{PI} \frac{g_i}{g_j} \frac{\hbar^2 \omega^2}{4\pi^2 m^2 c^2 v^2}
\]

where \( v \) is the velocity of the photoelectron. The recombination rate coefficient is obtained from the Maxwellian average as

\[
a_{RC}(T) = \frac{4}{\sqrt{\pi}} \left( \frac{m}{2kT} \right)^{3/2} \int_0^\infty v^3 e^{-\frac{m v^2}{2kT}} \sigma_{RC} dv,
\]

The recombination rate coefficient in terms of photoelectron energy can be obtained as \( a_{RC}(E) = \omega \sigma_{RC} \). Contributions from levels \( n_o \leq n \leq \infty \) can be obtained from the extension of Bell and Seaton theory [24] as described in [17,18]. Radiation damping [25] is included for fine structure levels.

### 2.4. Electron Impact Excitation (EIE)

The scattering matrix \( S_{SL\pi}(i,j) \) for excitation from \( i \) to \( j \) is obtained from the phase shift of the wave function which gives the EIE collision strength

\[
\Omega(S_i L_l - S_j L_{l_j}) = \frac{1}{2} \sum_{SL_{\pi} l_{l_j}} (2S + 1)(2L + 1) |S_{SL\pi}(S_i L_l - S_j L_{l_j})|^2,
\]

where SL denotes the symmetry of a state, initial or final, with S for the spin multiplicity although conventionally expressed only by S in a symmetry notation and L for the total angular momentum. S with subscripts \( SL\pi \) denotes the scattering matrix, e.g., \( S_{SL\pi}(S_i L_l - S_j L_{l_j}) \) is the scattering matrix for excitation between the two states, \( S_i L_l - S_j L_{l_j} \) with partial waves \( l_i \) and \( l_j \). \( \Omega \) is related to scattering cross-section \( \sigma_{EIE} \) as, \( \sigma_{EIE} = \pi/(g_i k^2) \Omega_{SR} \) where \( \pi \) is the standard constant of value 3.14,
$k$ is the momentum of the projectile and $a_0$ is the Bohr radius. The effective collision strength $\Upsilon(T)$ and excitation rate coefficient $q_{ij}(T)$ needed for astrophysical models are obtained as

$$Y(T) = \int_0^\infty \Omega_{ij}(\epsilon) \epsilon e^{-\epsilon/kT} d(\epsilon/kT), \quad q_{ij}(T) = \frac{8.63 \times 10^{-6}}{8_2 T^{1/2}} e^{-E_{ij}/kT} Y(T) \text{cm}^3\text{s}^{-1} \quad (22)$$

where $E_{ij} = E_j - E_i$ is in Ry, $T$ is in K, i.e., $1/kT = 157885/T$.

### 3. Data Description of NORAD-Atomic-Data

The NORAD-Atomic-Data webpage [1] contains general information on the data, such as description of the names of the files and notation, accuracy information, and data tables. It has the two data tables, one with nuclear charge $Z$ lower and the other with higher than 28. The elements with $Z \leq 28$ are H, He, C, N, O, F, Ne, Na, Mg, Al, Si, P, S, Cl, Ar, Ca, Ti, Cr, Fe and Ni. Since data are added with publications, each element has results for one, but typically more, ions and one or more atomic processes of photoexcitation, photoionization, electron–ion recombination, and electron impact excitation. As explained above, each process corresponds to the number of data files. The elements with $Z > 28$ are Br, W, Pt, Au, Pb, U. The atomic data for them contain energies and various types of transitions (E1, E2, E3, M1, M2) of their ions. Files containing Kα and Kβ transitions and lifetimes for levels of these ions, obtained under publications, are expected to be added.

Each row in the atomic data Tables 1 and 2 specifies the ion spectrum name and provides files for various atomic processes along the columns as shown in Figure 3. The names of the files are defined on the web page. Each atomic data file starts with the reference, the atomic process, contents of the file, description of data, and table of data. Some sample data files are described below.

**Figure 3.** Screenshot of the table that gives access to various atomic data. The leftmost column gives the atom/ion name and the right columns give the names of the data files for various atomic processes.

The tables of calculated energies contain $LS$ states or fine structure levels with spectroscopic identification. The identification is carried out after computation by the R-matrix method. The formats of the tables of energies obtained from the R-matrix method, either in $LS$ coupling or BPRM method, are different from those obtained from SUPERSTRUCTURE. SS assigns a single spectroscopic
designation for a level that corresponds to the highest mixing coefficient. Energies in $LS$ coupling approximation obtained from the R-matrix method are also uniquely identified with a single designation but in a different format. These energies are identified based on maximum channel percentage contribution and quantum defects. However, a BPRM level has one or more than one possible designation since contributions to the mixing coefficients outside the R-matrix boundary are not determined.

Table 1. Sample table of fine structure energy levels of Fe XIV [26] where levels have been grouped together as components of $LS$ terms. This is a conventional format of the NIST table [27].

<table>
<thead>
<tr>
<th>$C_t(S_tL_t\pi_t)$</th>
<th>$J_t$</th>
<th>$nl$</th>
<th>$2J$</th>
<th>$E(\text{Ry})$</th>
<th>$v$</th>
<th>$SL\pi$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nlv = 2, $^2L'': P\ (3\ 1)/2$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2p6s2</td>
<td>(1Se)</td>
<td>0</td>
<td>3p</td>
<td>1</td>
<td>$-2.88230 \times 10^1$</td>
<td>2.64</td>
</tr>
<tr>
<td>2p6s2</td>
<td>(1Se)</td>
<td>0</td>
<td>3p</td>
<td>3</td>
<td>$-2.86520 \times 10^1$</td>
<td>2.62</td>
</tr>
<tr>
<td>Nlv(c) = 2 : set complete</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Equiv electron/unidentified levels, parity: $e$

| $3s3p2$ | | 1 | $-2.68030 \times 10^1$ | 2.70 | 4P e |
| $3s3p2$ | | 3 | $-2.67330 \times 10^1$ | 2.71 | 4P e |
| $3s3p2$ | | 5 | $-2.66410 \times 10^1$ | 2.71 | 4P e |
| Nlv(c) = 3 : set complete | | | | | | |

| Nlv = 9, $^2L'': S\ (1)/2 P\ (3\ 1)/2 D\ (5\ 3)/2 F\ (7\ 5)/2 G\ (9\ 7)/2$ | | | | | | |
| $3p2$ | (1De) | 2 | 3d | 5 | $-1.96549 \times 10^1$ | 2.84 | 2DF e |
| $3p2$ | (1De) | 2 | 3d | 7 | $-1.95955 \times 10^1$ | 2.83 | 2FG e |
| $3p2$ | (1De) | 2 | 3d | 7 | $-1.94588 \times 10^1$ | 2.85 | 2FG e |
| $3p2$ | (1De) | 2 | 3d | 9 | $-1.94215 \times 10^1$ | 2.84 | 2G e |
| $3p2$ | (1De) | 2 | 3d | 3 | $-1.94120 \times 10^1$ | 2.83 | 2D e |
| $3p2$ | (1De) | 2 | 3d | 5 | $-1.93740 \times 10^1$ | 2.85 | 2D e |
| $3p2$ | (1De) | 2 | 3d | 1 | $-1.88526 \times 10^1$ | 2.85 | 2SP e |
| $3p2$ | (1De) | 2 | 3d | 1 | $-1.87559 \times 10^1$ | 2.86 | 2SP e |
| $3p2$ | (1De) | 2 | 3d | 3 | $-1.87283 \times 10^1$ | 2.88 | 2PD e |
| Nlv(c) = 9 : set complete | | | | | | |

Table 1 is a table of energies for Fe XIV that have been obtained using the BPRM method [26] and gives an example of the energy format. The levels have been identified with spectroscopic designation and are grouped together as fine structure levels of an $LS$ term. In addition to this format, the energy file also contains the same set of energies but listed in $J\pi$ order for modeling purposes. All notations in the file are in ASCII.

Tables of transitions, photoionization from the BPRM method and electron–ion recombination from the unified method contain numerical data for direct applications. The files have references at the top and a description of data below it. The initial and final levels of transitions, and levels for which photoionization cross-sections have been obtained from the R-matrix calculations and recombination rates and cross-sections from the unified method are encoded with numbers. These numerical identifications can be easily decoded, following the notes in the file and also matching the number with that in the energy file. A couple of samples of transition data files are given below. A few illustrations of use of photoionization data are given in the next Accuracy section. Table 2 illustrates the numerical format of transitions where transitions in Fe XIV have been chosen. Compared to this numerical format, the SS tables of transitions are more descriptive spectroscopically.
Table 2. Sample set of $f$-, $S$ and $A$-values for allowed E1 transitions in Fe XIV [26].

<table>
<thead>
<tr>
<th>$I_i$</th>
<th>$I_k$</th>
<th>$\lambda$(Å)</th>
<th>$E_i$(Ry)</th>
<th>$E_k$(Ry)</th>
<th>$f$</th>
<th>$S$</th>
<th>$A_k$(s$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>0</td>
<td>2</td>
<td>79</td>
<td>82</td>
<td>6478</td>
<td>$g_i$</td>
<td>$g_k$ $P_i$ $P_k$ $N_i$ $N_k$ $N_N$</td>
</tr>
<tr>
<td>1 1</td>
<td>451.12</td>
<td>$-2.6803\times10^1$</td>
<td>$-5.7777\times10^{-4}$</td>
<td>$1.716\times10^{-3}$</td>
<td>$1.893\times10^7$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 2</td>
<td>237.74</td>
<td>$-2.6803\times10^1$</td>
<td>$-1.231\times10^{-4}$</td>
<td>$1.927\times10^{-4}$</td>
<td>$1.453\times10^7$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 3</td>
<td>211.68</td>
<td>$-2.6803\times10^1$</td>
<td>$-2.819\times10^{-1}$</td>
<td>$3.929\times10^{-1}$</td>
<td>$4.197\times10^7$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 4</td>
<td>207.44</td>
<td>$-2.6803\times10^1$</td>
<td>$-1.458\times10^{-3}$</td>
<td>$1.991\times10^{-3}$</td>
<td>$2.259\times10^8$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 5</td>
<td>161.86</td>
<td>$-2.6803\times10^1$</td>
<td>$-4.713\times10^{-4}$</td>
<td>$5.023\times10^{-4}$</td>
<td>$1.200\times10^8$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 6</td>
<td>19.07</td>
<td>$-2.6803\times10^1$</td>
<td>$-4.846\times10^{-7}$</td>
<td>$6.086\times10^{-8}$</td>
<td>$8.890\times10^6$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 7</td>
<td>82.85</td>
<td>$-2.6803\times10^1$</td>
<td>$-5.076\times10^{-5}$</td>
<td>$2.769\times10^{-5}$</td>
<td>$4.931\times10^7$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 8</td>
<td>82.65</td>
<td>$-2.6803\times10^1$</td>
<td>$-1.231\times10^{-5}$</td>
<td>$6.699\times10^{-6}$</td>
<td>$1.202\times10^7$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 9</td>
<td>81.13</td>
<td>$-2.6803\times10^1$</td>
<td>$-1.5777\times10^{-5}$</td>
<td>$9.386\times10^{-6}$</td>
<td>$1.780\times10^7$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 10</td>
<td>79.57</td>
<td>$-2.6803\times10^1$</td>
<td>$-1.5351\times10^{-5}$</td>
<td>$1.716\times10^{-5}$</td>
<td>$8.989\times10^{-6}$</td>
<td>$1.807\times10^7$</td>
<td></td>
</tr>
</tbody>
</table>

NORAD-Atomic-Data also contains an additional file containing a set of transitions obtained from the BPRM method but with full spectroscopic designation (hence no need to use the energy file) for diagnostics of lines. This set corresponds to transitions among levels that have been observed, for example, the available observed energy levels from the NIST website [27]. Figure 4 gives a screenshot of a sample format of such a table. It presents fine-structure transitions in Fe XV with spectroscopic notation [28] as an example of transitions. Similar to the conventional format in the NIST table, fine structure transitions here have been grouped together as components of a $LS$ multiplet.

---


Fe XV: Oscillator strengths $f$, $S$, $A$-values for allowed E1 fine structure transitions with the observed energy level only.

Process: Fe XV $\rightarrow$ Fe XIV $ightarrow$ Fe XIII

File contents:

Table of fine-structure transitions among observed levels - (Table II in the paper).

$N$ = 26, $N_s$ = 11, $N_{ls}$ = 66

Number of observed levels = 66

No of transitions = 1030

Table:

<table>
<thead>
<tr>
<th>$C_i$</th>
<th>$C_j$</th>
<th>$f_{ls}$</th>
<th>$s_{ls}$</th>
<th>$g_{Is}$</th>
<th>$g_{Ij}$</th>
<th>$E_{is}$</th>
<th>$E_{ij}$</th>
<th>$f$</th>
<th>$S$</th>
<th>$A_{ij}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>3p3s2</td>
<td>3s3p</td>
<td>3s 3p 1 1 3 1</td>
<td>417.27</td>
<td>2.36 $\times 10^{-7}$</td>
<td>4.4 $\times 10^{-7}$</td>
<td>4.16 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3p3s2</td>
<td>3p3p</td>
<td>3p 3p 1 3 4 2</td>
<td>584.17</td>
<td>7.98 $\times 10^{-7}$</td>
<td>7.54 $\times 10^{-7}$</td>
<td>7.21 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3p3p</td>
<td>3p3p</td>
<td>3p 3p 1 3 4 2</td>
<td>584.17</td>
<td>7.98 $\times 10^{-7}$</td>
<td>7.54 $\times 10^{-7}$</td>
<td>7.21 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3p3s2</td>
<td>3p3p</td>
<td>3s 3p 1 3 3 4</td>
<td>100.38</td>
<td>8.25 $\times 10^{-7}$</td>
<td>8.27 $\times 10^{-7}$</td>
<td>1.11 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3p3s2</td>
<td>3p3s</td>
<td>3s 3p 1 3 3 5</td>
<td>93.03</td>
<td>1.26 $\times 10^{-7}$</td>
<td>1.27 $\times 10^{-7}$</td>
<td>1.29 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3p3s2</td>
<td>3p3p</td>
<td>3p 3p 1 3 4 2</td>
<td>584.17</td>
<td>7.98 $\times 10^{-7}$</td>
<td>7.54 $\times 10^{-7}$</td>
<td>7.21 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3p3p</td>
<td>3p3p</td>
<td>3p 3p 1 3 1 2</td>
<td>38.96</td>
<td>6.11 $\times 10^{-7}$</td>
<td>5.84 $\times 10^{-7}$</td>
<td>5.80 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3s3p</td>
<td>3p3p</td>
<td>3s 3p 3 1 1 2</td>
<td>317.59</td>
<td>5.16 $\times 10^{-7}$</td>
<td>2.74 $\times 10^{-7}$</td>
<td>1.73 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3s3p</td>
<td>3p3p</td>
<td>3s 3p 3 1 1 2</td>
<td>317.59</td>
<td>5.16 $\times 10^{-7}$</td>
<td>2.74 $\times 10^{-7}$</td>
<td>1.73 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3s3p</td>
<td>3p3p</td>
<td>3s 3p 3 1 1 2</td>
<td>317.59</td>
<td>5.16 $\times 10^{-7}$</td>
<td>2.74 $\times 10^{-7}$</td>
<td>1.73 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3s3p</td>
<td>3p3p</td>
<td>3s 3p 3 1 1 2</td>
<td>317.59</td>
<td>5.16 $\times 10^{-7}$</td>
<td>2.74 $\times 10^{-7}$</td>
<td>1.73 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>3s3p</td>
<td>3p3p</td>
<td>3s 3p 3 1 1 2</td>
<td>317.59</td>
<td>5.16 $\times 10^{-7}$</td>
<td>2.74 $\times 10^{-7}$</td>
<td>1.73 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3s3p</td>
<td>3p3p</td>
<td>3s 3p 3 1 1 2</td>
<td>317.59</td>
<td>5.16 $\times 10^{-7}$</td>
<td>2.74 $\times 10^{-7}$</td>
<td>1.73 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3s3p</td>
<td>3p3p</td>
<td>3s 3p 3 1 1 2</td>
<td>317.59</td>
<td>5.16 $\times 10^{-7}$</td>
<td>2.74 $\times 10^{-7}$</td>
<td>1.73 $\times 10^{-7}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 4. Screenshot of a transition table, with transitions in Fe XV as example [28], with full spectroscopic designation.
References of papers [29–111], which reported the atomic data of various processes in publications are added at end of the references which have been referred to in the text.

4. Accuracy of Data and Benchmarking

As mentioned, the atomic data at NORAD-Atomic-Data correspond to use of the R-matrix method with the close coupling approximation and atomic structure calculations with the Thomas–Fermi–Dirac–Amaldi potential for wave functions and energies. The results have been obtained in several approximations, such as LS coupling, fine structure splitting using algebraic transformation, relativistic Breit–Pauli approximation.

The R-matrix method with the CC approximation is known to be the most powerful approach to study atomic processes. The accuracy of atomic data computed depends on optimizations of a number of factors, such as (i) accuracy on the target or core ion wave functions which initiate the R-matrix computations, (ii) number of core ion excitations included in the wave function expansion, (iii) inclusion of relevant configurations in the bound-channel expansion (the second term of the close-coupling wave function expansion) that contribute to the total wave function expansion, (iv) the size of the R-matrix basis set representing the wave function in the inner region of the R-matrix sphere and (v) the size of the R-matrix sphere. During the study of an atomic process for an ion, a best possible balanced condition with these optimizations, computational power at OSC and time required for completion of computation is reached. Hence, accuracy of the atomic data is ion-specific, that is, it depends on the accuracy of the wave function representing the ion for the atomic process.

The accuracy for the Thomas–Fermi–Dirac–Amaldi potential for atomic structure calculations depends on the number of configurations and scaling of the orbital wave functions. SS can handle fewer of configurations than what R-matrix codes can do. R-matrix codes can get transitions, energies up to n = 10. However, the accuracy of SS can reach to that of program GRASP [112] in the Dirac–Hartree–Fock approximation for many cases, but SS is more robust than other higher accuracy codes in computing data with consistent accuracy except for very weak transitions.

For the extensive amount of work carried out under the OP and IP, the accuracy of data varied from less than 1% to 15% for most cases, but some data had larger differences from the existing accurate data in the literature. Based on the ab initio nature and consideration of a large number of configurations, and inclusion of relativistic effects, the accuracy for the R-matrix results, depending on the ion, can be estimated as follows:

(i) from less than 1 to 20% in LS coupling approximation,
(ii) from less than 1 to 50% for fine-structure splitting of LS multiplets using algebraic transformation,
(iii) from less than 1 to 15% for relativistic Breit–Pauli approximations.

For SS results in the Breit–Pauli approximation, the accuracy has been found to vary between less than 1% and 30–40% for transition rates.

One crucial test of accuracy for theoretical results is done through benchmarking with experimental measurements. Theoretical photoionization cross sections at NORAD-Atomic-Data have been benchmarked with measured spectra of photoionization at different experimental setups as follows:

1. with Advanced Light Source (ALS) at Lawrence Berkeley National Lab [113–117],
2. with Aarhus University set-up in Denmark [118,119],
3. with Orsay Lab at University De Paris-SUD set up in France [120,121],
4. with BESSY II set-up in Germany [122,123],
5. with pulsed-laser excitation and ionization atoms in an atomic beam at University of Nebraska [124].

Recombination cross-sections from unified method at NORAD-Atomic-Data have been benchmarked with measured spectra at experimental setups as follows:
Most users of NORAD-Atomic-Data are astronomers, physicists, and engineers. A few examples of applications are given below:

1. **Physics:** Photoionization cross-sections from NORAD-Atomic-Data were used to benchmark the cross-sections of N IV measured at the synchrotron facility BESSY II setup by Simon et al. [122] (Figure 5).

2. **Astronomy:** Figure 6 shows that the use of detailed photoionization cross sections [127] available at NORAD for Cr-to-Fe ratio as a probe for chemical evolution between NLTE analysis of Cr I and Cr II lines resulted in good agreement [128].

3. **Engineering:** Adeneh et al. [129] who studied thermodynamic and radiative properties of electrical discharge machining (EDM) plasmas for temperature up to 10,000 K and pressure range 0.1–1 MPa using atomic data from NORAD-Atomic-Data, find an increase in net emission coefficient (NEC) with different amount of iron contamination in nitrogen and sharp cooling of the plasma by iron contamination as shown in Figure 7 taken from [129].

**Figure 5.** Comparison of photoionization cross sections of N IV measured at BESSY II setup in Germany by Simon et al. [122] (upper panel) with those by Nahar and Pradhan [123] available at the NORAD-Atomic-Data (blue) and MCDF calculations (orange drop [122]) in the lower panel (Figure from [122] is used with permission).

**Figure 6.** Cr-to-Fe ratio as probe of chemical evolution between NLTE analysis of Cr I and Cr II lines (left panel) obtained by using photoionization cross sections at NORAD (right panel [127]) [128]. (Figure from [128] is used with permission).
5. Conclusions

An overview on the contents, theoretical outline for the methods used for the data, accuracy and benchmarking of the online database, NORAD-Atomic-Data, is presented. Future directions are:

(i) include more data with publications,
(ii) include computer programs that can read the data files and process the data to calculate the quantities of interest,
(iii) introduce a plotting feature,
(iv) an option for selection of partial data, such as, for a particular wavelength range.

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Conflicts of Interest: The author declares no conflict of interest.

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