



Article The XSTAR Atomic Database

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Abstract: We describe the atomic database of the XSTAR spectral modeling code, summarizing the systematic upgrades carried out in the past twenty years to enable the modeling of K-lines from chemical elements with atomic number $Z \leq 30$ and recent extensions to handle high-density plasmas. Such plasma environments are found, for instance, in the inner region of accretion disks round compact objects (neutron stars and black holes), which emit rich information about the system's physical properties. Our intention is to offer a reliable modeling tool to take advantage of the outstanding spectral capabilities of the new generation of X-ray space telescopes (e.g., XRISM and ATHENA) to be launched in the coming years. Data curatorial aspects are discussed and an updated list of reference sources is compiled to improve the database provenance metadata. Two XSTAR spin-offs—the ISMabs absorption model and the uaDB database—are also described.

Keywords: XSTAR; atomic databases; atomic processes; line formation; X-rays; high-density plasmas

1. Introduction

The XSTAR (https://heasarc.gsfc.nasa.gov/docs/software/xstar/xstar.html) code computes the physical conditions and emission spectra of a photoionized gas and has been widely used in astrophysics, most notably in X-ray astronomy, for the past 20 years [1,2]. It assumes an ionizing radiation source surrounded by a spherical gas shell that absorbs and transfers parts of this radiation to finally emit a spectrum. From an input comprising the incident continuum, shell thickness, elemental abundances, and gas density, the code computes the ionization balance, level populations, and temperature to generate the gas opacity and the emitted line and continuum fluxes. The computational model determines simultaneously the gas state and radiation field at every user-defined spatial zone of the cloud. To ensure convergence in local thermodynamic equilibrium (LTE) conditions, it implements a detailed and consistent treatment of the radiative (bound–bound, bound–free, and free–free) and collisional (electron-impact excitation and ionization, electron–ion recombination, charge transfer and three-body recombination) processes, a procedure that in turn depends on the accuracy of the underlying database of atomic rate coefficients.



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Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https://creativecommons.org/licenses/by/4.0/). Therefore, XSTAR is an ideal platform to study data curation schemes for implementing and maintaining application-based atomic databases.

Since 2001, we have been systematically computing for the XSTAR database the atomic data required to model the K-lines and edges of both cosmically abundant and trace elements [3–46]. These X-ray spectral features are prime candidates to devise diagnostics to determine the plasma conditions and chemical abundances in a wide variety of astronomical entities: interstellar, intergalactic, and intra-cluster media; active galactic nuclei (AGN); X-ray binaries; supernova remnants; stars; the Sun, and some solar planets and satellites. This long-term collaboration has reached an important milestone, and the corresponding upgrading of the XSTAR database with the new datasets has more than doubled its volume, bringing to the fore a series of data curational problems discussed in this report. Our final goal is to upgrade XSTAR to take advantage of the relatively high spectral resolution (5 eV in the soft X-ray band \sim 0.3–12 keV) and sensitivity of the microcalorimeter-based spectrometers in the joint NASA/JAXA X-ray Imaging and Spectroscopy Mission (XRISM, https://heasarc.gsfc.nasa.gov/docs/xrism) to be launched in 2022 and in the European Space Agency flagship Advanced Telescope for High-ENergy Astrophysics (ATHENA, https://sci.esa.int/web/athena). As an example of this accuracy level, Figure 1 shows the XSTAR fit in the 6.50–6.60 keV region of the spectrum of the Perseus Cluster taken by the HITOMI (https://www.nasa.gov/hitomi) space telescope before its demise in 2016 [47].



Figure 1. HITOMI spectrum of the Perseus Cluster in the Fe XXV "triplet" spectral region (6.50–6.60 keV) [47] showing the XSTAR fit (red line). With the higher resolution, the He-like "triplet" becomes a quartet comprising the resonance (w), magnetic quadrupole (x), intercombination (y), and magnetic dipole (z) lines. Other Fe satellite lines begin to be detectable (e.g., the bump at ~6.5 keV).

We have recently introduced high-density ($n_e > 10^{18} \text{ cm}^{-3}$) effects in the XSTAR database, which come into play in the reflection spectra of the inner region of the accretion disks around compact objects (e.g., black-hole candidates and neutron stars): continuum lowering [40–45]; dielectronic recombination suppression [48,49]; collisional ionization; three-body recombination; and stimulated emission. Since the Fe abundance is a key measure of line reprocessing in reflection models [50], we are now in a good position to test if the neglect of such high-density effects is responsible for the anomalously high abundances recurrently derived [51]. Regarding XSTAR database curation, the inclusion of

high-density effects presents serious execution difficulties that are now managed by preprocessing the database at each density point. Since this pre-processing time is generally long, a grid of pre-computed databases has been deployed that, considering the data volumes involved, favors a cloud computing environment for the XSTAR modeling of high-density plasmas.

Finally, the development of the XSTAR database has given rise to two spin-offs: the ISMabs (https://heasarc.gsfc.nasa.gov/xanadu/xspec/models/ismabs.html) absorption model [35] and the uaDB (https://heasarc.gsfc.nasa.gov/uadb/index.php) online database. The former is the result of benchmarks of the theoretical photoionization curves with observed K absorption in the interstellar medium (ISM), which has led to the implementation of a reliable model of ISM K absorption and to the matching of the theoretical K-line wavelengths with the astronomically observed and laboratory measured values, the latter two not always agreeing (e.g., O I [23,52]). The uaDB database allows the downloading of the XSTAR and other atomic rates without having to interfere with the code.

2. XSTAR Atomic Database

The XSTAR database comprises records of multilevel model ions (usually referred to as ion targets) and data structures to derive the rates for the microprocesses that determine the ionization and excitation of the plasma [2]. The rate types (*rt*) are the following:

- **01.** Ground state ionization
- **03.** Bound–bound collision
- 04. Bound-bound radiative
- 05. Bound–free collision (level)
- **06.** Total recombination
- 07. Bound–free radiative (level)
- 08. Total recombination forcing normalization
- **09.** Two-photon decay
- **11.** Element data
- 12. Ion data
- **13.** Level data
- 14. Bound–bound radiative, superlevel to spectroscopic level
- 15. Collisional ionization total rate
- 40. Bound-bound collisional, superlevel to spectroscopic level
- **41.** Non-radiative Auger transition
- 42. Inner-shell photoabsorption followed by autoionization.

For each ionic species (Z; N) in the ranges $1 \le Z \le 30$ and $1 \le N \le Z$, where *Z* and *N* are, respectively, the atomic and electron numbers, the data for the target models and processes are contained in ASCII files labeled d*ion*t0*dt*. The ion index *ion* is defined as

$$ion = Z(Z-1)/2 + (Z-N) + 1,$$
(1)

and dt is the data type (see Appendix A for a complete list). Each diont0dt file contains integer, floating-point, and character variables. For each release, the database is transcribed to a binary FITS-format-derived [53,54] structure (atdb.fits) consisting of three onedimensional arrays (floating point, integer, and character) and a pointer array, which are uploaded in main memory at runtime. This scheme ensures portability, fast data uploading when the code is invoked, and single-data accesses in main memory during processing.

The XSTAR data curation strategy essentially follows two principles:

1. Since the datasets are compiled from diverse sources, the original data structures and units are explicitly kept in the database records to be transcribed and unified at runtime. For instance, level energies are specified in eV but the photoionization cross-sections are usually tabulated in Rydbergs. Levels in some atomic models are listed in intermediate coupling (fine structure) while in others *LS* terms are adopted.

These format differences are accounted for by defining data types, each of which adheres to its own set of rules for data tabulation.

2. As new data and processes are included in the database, new data types may be defined. Thus, photoionization cross-sections can be specified in more than one data type—valence shell and K shell, say—if the tabulation for each energy region comes from different sources. They are then concatenated and adjusted at runtime.

We show in Figure 2 the target model for Mg-like Zn XIX complying with data type dt = 06 and contained in file d454t006. Each row is headed by the integer tuple $(dt, rt, 0, N_{\text{flt}}, N_{\text{int}}, N_{\text{chr}})$, where dt is the data type, rt the rate type, N_{flt} the number of floating-point parameters, N_{int} number of integer parameters, and N_{chr} the length of the character string. As shown in the rate-type list above, rt = 13 corresponds to level data. The first and second floating-point columns tabulate level energies (eV) and statistical weights g = 2J + 1 (the other two floating-point columns have been edited out). The second integer tuple (n, 2S + 1, L, Z, i, ion) lists: the *n* principal quantum number of the valence electron; the 2S + 1 spin multiplicity; the total orbital angular momentum *L*; element atomic number *Z*; the *i* level index, and *ion* index. The character string gives the level assignment (configuration and spectroscopic term), the end of the record being indicated with the % character. Records may span one arbitrary length line or multiple lines in the ASCII file.

6	13	0	4	6	8	0.000000E+00	1.000000E+00 .	 3	1	0	30	1	454	3s2.1S_0 %
6	13	0	4	6	12	3.6651995E+01	1.000000E+00 .	 3	3	1	30	2	454	3s1.3p1.3P_0 %
6	13	0	4	6	12	3.8015000E+01	3.000000E+00 .	 3	3	1	30	3	454	3s1.3p1.3P_1 %
6	13	0	4	6	12	4.1638999E+01	5.000000E+00 .	 3	3	1	30	4	454	3s1.3p1.3P_2 %
6	13	0	4	6	12	5.6705005E+01	3.000000E+00 .	 3	1	1	30	5	454	3s1.3p1.1P_1 %
6	13	0	4	6	12	8.8177000E+01	1.000000E+00 .	 3	3	1	306	5 45	54 35	s0.3p2.3P_0 %
6	13	0	4	6	12	8.9971999E+01	3.000000E+00 .	 3	3	1	30	7	454	3s0.3p2.3P_1 %
6	13	0	4	6	12	9.0909009E+01	5.000000E+00 .	 3	3	1	30	8	454	3s0.3p2.3P_2 %
6	13	0	4	6	12	9.5230001E+01	5.000000E+00 .	 3	1	2	30	9	454	3s0.3p2.1D_2 %
6	13	0	4	6	12	1.0956001E+02	1.000000E+00 .	 3	1	0	30	10	454	3s0.3p2.1S_0 %
6	13	0	4	6	16	1.0583000E+03	3.000000E+00 .	 3	3	0	30	11	454	2p5.3s2.3p1.3S_1 %
6	13	0	4	6	16	1.0607000E+03	5.000000E+00 .	 3	3	2	30	12	454	2p5.3s2.3p1.3D_2 %
6	13	0	4	6	16	1.0645000E+03	7.000000E+00 .	 3	3	2	30	13	454	2p5.3s2.3p1.3D_3 %
6	13	0	4	6	16	1.0654000E+03	3.000000E+00 .	 3	1	1	30	14	454	2p5.3s2.3p1.1P_1 %
6	13	0	4	6	16	1.0674000E+03	5.000000E+00 .	 3	3	1	30	15	454	2p5.3s2.3p1.3P_2 %
6	13	0	4	6	16	1.0766000E+03	1.000000E+00 .	 3	3	1	30	16	454	2p5.3s2.3p1.3P_0 %
6	13	0	4	6	16	1.0843000E+03	3.000000E+00 .	 3	3	2	30	17	454	2p5.3s2.3p1.3D_1 %
6	13	0	4	6	16	1.0895000E+03	3.000000E+00 .	 3	3	1	30	18	454	2p5.3s2.3p1.3P_1 %
6	13	0	4	6	16	1.0901000E+03	5.000000E+00 .	 3	1	2	30	19	454	2p5.3s2.3p1.1D_2 %
6	13	0	4	6	16	1.1029000E+03	1.000000E+00 .	 3	1	0	30	20	454	2p5.3s2.3p1.1S_0 %
6	13	0	4	6	20	1.2384461E+03	1.000000E+00 .	 3	3	1	30	21	454	2s1.2p6.3s2.3p1.3P_0 %
6	13	0	4	6	20	1.2391314E+03	3.000000E+00 .	 3	3	1	30	22	454	2s1.2p6.3s2.3p1.3P_1 %
6	13	0	4	6	20	1.2434606E+03	5.000000E+00 .	 3	3	1	30	23	454	2s1.2p6.3s2.3p1.3P_2 %
6	13	0	4	6	20	1.2457835E+03	3.000000E+00 .	 3	1	1	30	24	454	2s1.2p6.3s2.3p1.1P_1 %
6	13	0	4	6	24	9.7285003E+03	1.000000E+00 .	 3	3	1	30	25	454	1s1.2s2.2p6.3s2.3p1.3P_0 %
6	13	0	4	6	24	9.7293003E+03	3.000000E+00 .	 3	3	1	30	26	454	1s1.2s2.2p6.3s2.3p1.3P_1 %
6	13	0	4	6	24	9.7340004E+03	5.000000E+00 .	 3	3	1	30	27	454	1s1.2s2.2p6.3s2.3p1.3P_2 %
6	13	0	4	6	24	9.7373003E+03	3.000000E+00 .	 3	1	1	30	28	454	1s1.2s2.2p6.3s2.3p1.1P_1 %
6	13	0	4	6	10	6.6340128E+02	2.000000E+00 .	 12	1	0	30	29	454	superlevel %
6	13	0	4	6	14	1.0768748E+04	1.000000E+00 .	 1	0	0	30	30	454	<pre>superlevel_[K] %</pre>
6	13	0	4	6	9	6.9751000E+02	2.000000E+00 .	 3	2	0	30	31	454	continuum %

Figure 2. Content of file d454t006 of the XSTAR database listing the atomic model (data type dt = 06 and rate type rt = 13) for Mg-like Zn XIX (*ion* = 454).

This ion representation presents remarkable features. By looking at the level-*i* index and assignment, the Zn XIX model contains data attributes for 31 levels from: the valence shell $(1 \le i \le 10)$, open L shell $(11 \le i \le 24)$, and open K shell $(25 \le i \le 28)$; two superlevels (*i* = 29–30); and a continuum level (*i* = 31). The latter is associated to the ionization potential of the ion. Therefore, the model comprises spectroscopic bound levels, quasi-bound L- and K-vacancy resonances, and artificial levels. The tabulation must identify the ground level with index 1 and the continuum level with the highest index. Apart from this constraint, the levels are not necessarily listed in energy order facilitating model upgrading, which simply involves adding new levels at the end of the existing tabulation. The level-assignment string has been carefully devised and used throughout to facilitate searches based on filled shells, shell holes, *LS* terms, and *J* total angular momenta.

Following [1], the superlevel accounts in H- and He-like ions for the recombination cascade from unaccounted high-lying levels to the spectroscopic levels of the model atom. Rates have been determined from population kinetic calculations. For other isoelectronic sequences, the superlevel is assumed to decay radiatively and collisionally to the ground state. Rates populating and depopulating the superlevel are then calculated by fitting to the total recombination rate, thus explicitly accounting for the contributions from excitation, ionization, and recombination to the level populations. As a result, every ionic process is accompanied by its detailed-balance inverse, ensuring the progressive convergence to the LTE level populations. The superlevel_[K] is defined in Section 3.1.

All other data types follow similar data structures (floating point, integer, and character) as those displayed in Figure 2. For instance, we show in Figure 3 an excerpt of file d454t050 containing radiative data for the bound-bound transitions (upper level $k \le 10$) in Zn XIX. As before, the header integer tuple prescribes the data and rate types dt = 50 and rt = 4; three floating-point items; four integer items; and no character strings. The floating-point tuple $(\lambda, gf(i, k), A(k, i))$ lists the transition wavelength (Å), oscillator strength (gf-value), and transition probability (A-value, s^{-1}). The second integer tuple (*i*, *k*, *Z*, *ion*) lists the lower and upper levels of the transition, atomic number, and ion index. By comparing the *i* and *k* transition indices with the *i* level indices in Figure 2, it may be seen that the tabulation includes both allowed and forbidden transitions, e.g., the resonance transition (i,k) = (1,5) and the forbidden transition (i,k) = (1,4). Importantly, every excited level in the database must include a decay mechanism, radiative and/or collisional; otherwise, its population may grow out of bounds when modeling. For instance, the 3s1.3p1.3P_0 metastable level (i = 2) does not display a radiative transition ($j = 0 \rightarrow 0$ radiative transitions are strictly forbidden), and consequently, an alternative decay process must be provided in a separate data type; e.g., collisional de-excitation.

```
3.2047230E+02 5.2260000E-03 1.1310000E+08 1 3 30 454
50 4 0 3 4 0
                                                                     %
50 4 0 3 4 0
              2.0629320E+02 7.0930000E-01 3.7060000E+10 1 5 30 454
                                                                     %
50 4 0 3 4 0
              2.4417010E+02 2.5340000E-01 2.8350000E+10 3 6 30 454
                                                                     %
50 4 0 3 4 0
              4.2222080E+02 4.7650000E-03 1.7830000E+08 5 6 30 454
                                                                      %
50 4 0 3 4 0
              2.2582420E+02 2.7330000E-01 1.1910000E+10 2
                                                           7 30 454
                                                                      %
50 4 0 3 4 0
              2.3171380E+02 1.9730000E-01
                                           8.1690000E+09 3
                                                            7 30 454
                                                                      %
50 4 0 3 4 0
              2.4845700E+02 3.1050000E-01
                                           1.1180000E+10 4
                                                            7 30 454
                                                                      %
50 4 0 3 4 0
              3.8631020E+02 1.4800000E-03
                                           2.2060000E+07 5
                                                            7 30 454
                                                                      %
50 4 0 3 4 0
              2.2265070E+02 3.4070000E-01 9.1680000E+09 3 8 30 454
                                                                      %
              2.3806620E+02 7.8830000E-01 1.8560000E+10 4
50 4 0 3 4 0
                                                           8 30 454
                                                                      %
              3.6175990E+02 1.2440000E-01 1.2680000E+09 5 8 30 454
50 4 0 3 4 0
                                                                      %
50 4 0 3 4 0
              2,0084360E+02 2,0630000E-02 6,8230000E+08 3 9 30 454
                                                                     %
50 4 0 3 4 0
              2.1330280E+02 2.0500000E-01 6.0100000E+09 4 9 30 454
                                                                     %
50 4 0 3 4 0
              3.0751030E+02 8.4760000E-01 1.1960000E+10 5 9 30 454
                                                                     %
50 4 0 3 4 0
              1.7136220E+02 2.6040000E-03 5.9150000E+08 3 10 30 454
                                                                     %
50 4 0 3 4 0
              2.4339670E+02 3.8770000E-01 4.3650000E+10 5 10 30 454
                                                                     %
50 4 0 3 4 0
              2.9315000E+02 0.0000000E+00 9.5940000E+00 1 4 30 454
                                                                     %
```

Figure 3. Excerpt of file d454t050 of the XSTAR database listing the radiative transitions with upper level $k \le 10$ (data type dt = 50 and rate type rt = 4) in Mg-like Zn XIX (*ion* = 454).

Data type dt = 86, listing the radiative and Auger widths of K-vacancy levels, is also an interesting case as it involves transitions between different ion-charge states (Figure 4). The floating-point tuple $(E(k_N), A_a(k_N), A_a(k_N, i_{N-1}), A_r(k_N))$ lists: the energy $E(k_N)$ (in eV relative to the ionization threshold) and total Auger width $A_a(k_N)$ (s⁻¹) of the k_N level of the *N*-electron ion ($ion_N = 454$); the partial Auger width $A_a(k_N, i_{N-1})$ (s⁻¹) leaving the (N - 1)-electron ion ($ion_{N-1} = 455$) in the i_{N-1} level; and the total radiative width $A_r(k_N)$ (s⁻¹) of the k_N level of the *N*-electron ion. The second integer tuple then specifies ($i_{N-1}, k_N, Z, ion_{N-1}, ion_N$). In this example, the (N - 1)-electron ion is assumed to end up in its ground state ($i_{N-1} = 1$) after the Auger transition; however, this is not always the case as the Auger transition may involve an excited end level of the (N - 1)-electron ion. Such partial Auger widths are not determined in some structure codes (e.g., HFR [55]) and in the *R*-matrix scattering method [56,57], the latter including radiative and Auger damping through an optical model potential [58].

 86 4 0 4 5 0
 9.0309903E+03
 1.1100000E+15
 1.1100000E+15
 1.1600000E+15 1
 1 25 30 455 454 %

 86 4 0 4 5 0
 9.0317903E+03
 1.100000E+15
 1.100000E+15
 1.1800000E+15 1
 1 26 30 455 454 %

 86 4 0 4 5 0
 9.0364904E+03
 1.100000E+15
 1.100000E+15
 1.1600000E+15 1
 1 27 30 455 454 %

 86 4 0 4 5 0
 9.0397903E+03
 1.080000E+15
 1.080000E+15
 1.290000E+15 1
 28 30 455 454 %

Figure 4. Content of file d454t086 of the XSTAR database listing the radiative and Auger widths (data type dt = 86 and rate type rt = 4) of Mg-like Zn XIX (*ion* = 454).

Table A1 in Appendix B tabulates the number of levels for each ionic species (Z; N). It may be seen that most neutral and, in many cases, the singly ionized systems are not well represented, indicating that XSTAR is mainly destined to model photoionized plasmas for which such species have negligible fractions. The number of levels in the ion models is never above 1000 to enable practical runtimes when modeling (see Section 6.1), and the most relevant elements in the astronomical soft X-ray band—N, O, Ne, Fe, and Ni—are especially well treated.

As fully described in the original paper on the XSTAR database [2], atomic datasets have been collected from a myriad of individual sources and data compilations such as the NIST Atomic Spectra Database (https://www.nist.gov/pml/atomic-spectra-database) [59], TOPbase (http://cdsweb.u-strasbg.fr/topbase/topbase.html) [60], CHIANTI (https://www.chiantidatabase.org/) [61], ADAS (https://www.adas.ac.uk/), Iron Project (http://cdsweb.u-strasbg.fr/topbase/testop/TheIP.html) [62], and PP95 (https://ned.ipac.caltech. edu/level5/Pradhan/Pradhan_contents.html) [63]. Major upgrades have been mainly concerned with the radiative recombination (RR) and dielectronic recombination (DR) data from the AMDPP tabulations (http://amdpp.phys.strath.ac.uk/tamoc/DATA/) [64,65] and from the systematic computations of atomic data for K-line diagnostics summarized in the next Section 3. A complete data provenance list is given in Appendix C.

3. Computation of Atomic Data for K-Line Diagnostics

Since XSTAR is widely used for spectral modeling in X-ray astrophysics, we have dedicated a systematic effort to compute the atomic data to characterize K-lines in ionic species with $Z \leq 30$ —namely, valence and K-vacancy level energies, transition wavelengths, *A*-values, radiative and Auger widths, and photoionization cross-sections [5–29]. Due to the lack of spectroscopic data to validate most ionic models, a multi-code methodology was implemented right from the outset to compute the data and estimate their accuracy. Structure data were computed with the multiconfiguration codes HFR (Pauli Hamiltonian) [55], AUTOSTRUCTURE (Breit–Pauli Hamiltonian) [66,67], and GRASP92 (Dirac–Coulomb–Breit Hamiltonian) [68], and the photoionization cross-sections with the Breit–Pauli *R*-matrix suite of codes [56,57] and AUTOSTRUCTURE. Due to the large target models for species in isoelectronic sequences with electron number $19 \leq N \leq 26$, the latter code was used to compute their photoionization cross-sections in the distorted-wave approximation. Initial priority was given to ions from the oxygen, nitrogen, and iron isonuclear sequences and then from even-*Z* elements, but those from odd-*Z* and trace elements have been recently completed.

In the calculations of accurate atomic data for K-line diagnostics, two key effects must be considered in detail: radiative and Auger damping and orbital relaxation. In Sections 3.1 and 3.2, we briefly outline these two processes and the methods to include them in structure and scattering calculations.

3.1. Radiative and Auger Damping

Let us consider an ion, say, with electron number $N \le 10$. When a photon promotes a K-shell electron to an excited Rydberg state,

$$h\nu + 1s^2 2s^\lambda 2p^\mu \longrightarrow 1s 2s^\lambda 2p^\mu np$$
, (2)

it subsequently decays via the radiative and Auger manifold

$$1s2s^{\lambda}2p^{\mu}np \xrightarrow{Kn} 1s^{2}2s^{\lambda}2p^{\mu} + h\nu_{n}$$
(3)

$$\xrightarrow{\kappa\alpha} 1s^2 2s^{\lambda} 2p^{\mu-1} np + h\nu_{\alpha} \tag{4}$$

$$\overset{\text{KL}n}{\longrightarrow} \begin{cases} 1s^2 2s^{\lambda} 2p^{\mu-1} + e^- \\ 1s^2 2s^{\lambda-1} 2p^{\mu} + e^- \end{cases}$$

$$(5)$$

$$\stackrel{\text{KLL}}{\longrightarrow} \begin{cases} 1 s^2 2 s^{\lambda} 2 p^{\mu - 2} n p + e^- \\ 1 s^2 2 s^{\lambda - 1} 2 p^{\mu - 1} n p + e^- \\ 1 s^2 2 p^{\mu - 2} n p + e^- \end{cases}$$
(6)

dominated by K α radiative decay (Equation (4)) and KLL Auger spectator-electron ionization (Equation (6)). Such channeling causes a damping effect (see Figure 5) since the $1s2s^{\lambda}2p^{\mu}np$ resonances have broad and symmetric widths almost independent of n, which must be taken into account when devising plasma diagnostics. In the *R*-matrix method, it is computationally intractable to treat this channel array from first principles as it would involve target states with np orbitals in an ever increasing close-coupling expansion. Therefore, for the higher Rydberg resonances, damping is neatly accounted for via an additional imaginary potential $V_{\text{opt}} = -i(\Gamma_r + \Gamma_a)/2$ involving the total radiative width Γ_r —as developed in the Hickman–Robicheaux formalism [69,70] (which is essentially equivalent to the Davies–Seaton formalism [71,72])—and adapted to include the total Auger width Γ_a [58].



Figure 5. Photoabsorption cross-section of the 1s²2s²2p⁶ ¹S ground state of Ne-like Fe XVII in the K-edge region. *Top panel*: undamped cross-section. *Bottom panel*: damped (radiation and Auger) cross-section.

Due to their diagnostic importance, $K\alpha$ lines in the XSTAR database are treated as bound–bound transitions and assigned wavelengths and *A*-values, but the upper Kvacancy states also display Auger widths that lead to autoionization (see Figures 2–4). However, the K resonances also appear in the photoabsorption cross-sections, as shown in Figure 5, smearing the edge due to their peculiarly symmetric and constant line profiles. As a result, the K α resonances must be trimmed out from the partial photoionization cross-sections that are used to derive the rates so as not to count them twice. However, it is more complicated to carry out this trimming procedure in the L-edge region due to the piling up of a large number of resonances converging to closely spaced thresholds; thus some degree of double counting is practically unavoidable, making the L-edge less useful to devise plasma diagnostics.

As shown in Figure 6, damping leads to a significant difference between photoabsorption and photoionization, and in the XSTAR code, both are needed: the former to determine opacities and the latter to derive the photoionization rates. However, the database essentially stores partial photoionization cross-sections, i.e., cross-sections leaving the photoionized (N - 1)-electron system in specific states, and consequently, the difference between the total photoabsorption and photoionization cross-sections is assigned to superlevel_[K] (see level i = 30 in Figure 2).



Figure 6. K-edge photoabsorption (full curve) and photoionization (dotted curve) cross-sections of Ca XV (C-like) as a function of photon energy. The dashed curve gives the photoabsorption cross-section convolved with a Gaussian of 0.001 width. Reproduced from Figure 3 of [18] with permission of the AAS.

Due to the large number of target levels in ions with electron number N > 18, their photoionization cross-sections were computed using the simpler distorted-wave method implemented in AUTOSTRUCTURE [29]. The level numbers for such ionic systems tabulated in Table A1 (Appendix B) underwent considerable trimming to comply with the modeling specifications of XSTAR and, hence, do not represent the extent of the actual collisional targets. In AUTOSTRUCTURE, photoabsorption is conveniently treated as the sum of two separate processes: photoionization and photoexcitation. However, the complicated decay routes of K-vacancy states delineated in Equations (3)–(6) must be explicitly included in the photoexcitation process, and were thus computed in *LS* coupling. Since XSTAR requires fine-structure partial photoionization cross-sections to determine rates, we decided to obviate the phototexcitation process in ions with $Z \neq 26$ and $21 \leq N \leq 26$, thus underestimating their opacities, although the radiative and Auger decays of the L α , β and K α , β lines from these systems will be respectively taken into account in data types 50 and 86 (see Figures 3 and 4).

3.2. Orbital Relaxation

As shown in [55,73], the photoionization of an inner-shell electron leads to a *relaxation* of the outer electrons of the ionized state due to a reduced screening of the atomic nuclear potential by the reduced inner electron cloud; i.e., an increased effective charge Z_{eff} . For example, in the inner-shell photoionization of O I 1s²2s²2p⁴ to O II 1s2s²2p⁴, the effective charge experienced by a 2p electron in the initial and final states is a complicated average screening including the other equivalent 2p electrons. Nevertheless, by using hydrogenic considerations, it is found that the electron orbital energy ($E_n \sim -E_R Z_{\text{eff}}^2/n^2$) is more tightly bound for the relaxed 2p electron, and the electron density is concentrated closer to the nucleus ($r_n \sim a_0 n^2/Z_{\text{eff}}$).

To illustrate this phenomenon, we show in Figure 7 the outer 2p orbital in both neutral oxygen and its K-vacancy ionized daughter obtained from single-configuration Hartree–Fock calculations [74,75]. Clearly, the *relaxed* 2p orbital of the ionized O II inner-shell state differs significantly from the neutral O I 2p orbital, a difference that needs to be accounted for to compute more accurate atomic transition properties.



Figure 7. Probability density of the Hartree–Fock 2p orbital in the initial O I 1s²2s²2p⁴ ground state compared to the 2p orbital in the final O II 1s2s²2p⁴ K-vacancy state. The 2p electrons of the latter are screened from the nucleus by one fewer inner-shell 1s electron and are therefore "relaxed" to more tightly bound orbitals.

In our HFR and AUTOSTRUCTURE structure calculations, orbital relaxation was taken into account by adopting different non-orthogonal orbital bases for each configuration. In most photoionization calculations, however, an orthonormal orbital basis is used because of the relative simplicity in computing transition and energy matrix elements. Relaxation then requires a substantial multi-configurational wave-function description including additional orbitals to account for the significant differences between initial- and final-state orbitals. Thus, additional care must be taken in computing inner-shell photoionization cross-sections when compared to outer-shell photoionization, as discussed more fully in [23]. Due to these difficulties, most large-scale computations of photoabsorption and photoionization cross-sections do not include orbital relaxation effects unless otherwise indicated.

4. Photoabsorption Cross-Section Benchmarks

Table A2 in Appendix C lists the reference sources for the computations of the photoabsorption cross-sections. Measurement of K absorption in the laboratory and ISM enables useful benchmarks of the theoretical cross-sections to improve spectral accuracy. In the following sections, we give a summary of these procedures.

4.1. Carbon Sequence

Calculations of the atomic data to model the K-shell photoabsorption by carbon ions were performed by [21], and a benchmark was obtained using *Chandra* X-ray high-resolution spectra [37]. The C II K α "triplet" as well as the C III K α and K β lines were identified in the X-ray spectra of four super-soft sources. C I was not modeled due to the presence of instrumental features at ~43.6 Å.

4.2. Oxygen Sequence

K-shell photoabsorption cross-sections for O I–O VII were computed by [11], and comparisons of the theoretical resonance positions with astronomical observations and laboratory measurements were carried out by [32,33]. It was found that, due to a suspect experimental energy calibration scale, the X-ray astronomical observations provided a more reliable reference to perform the benchmark of the atomic data. For O I, the K-edge cross-section was re-examined in detail to yield a substantially improved version for the XSTAR atomic database [23].

In this exhaustive study, the K α (1s \rightarrow 2p) resonance in O I was examined with *R*matrix and Multi-Configuration Hartree-Fock (MCHF) calculations, existing laboratory spectroscopic data, and several X-ray observational assessments (see Table 1). While practical *R*-matrix calculations for multi-electron systems are better tailored for computing scattering and atomic transition rates, they are less reliable in predicting the energy differences between atomic states as compared to limited bound-state structure calculations. (R-matrix calculations necessarily include all quasi-bound and continuum final states, limiting a practical improvement of the basis description.) The *R*-matrix K α energy was found to be ~ 0.3 eV higher than the final assessed observational value of 527.37 eV [23]. More complex MCHF calculations, which were focused only on the initial $1s^22s^22p^4$ state and final 1s2s²2p⁵ state, converged downward to a value of 527.49 eV, in good agreement with the observations. On the other hand, the experiments performed at the Advanced Light Source (ALS) were reporting a value of 526.79 eV [76], a discrepancy of almost -0.6 eV. Considering the reliability of the observational calibration and the convergence trend of the MCHF calculations, the X-ray observation was unconventionally chosen instead of the laboratory measurement as the final resonance position in the atomic absorption model.

Method	Source	E(1s-2p)	E(1s-3p)	ΔE
Astronomical observations	XMM-Newton, Mrk 421 [23]	527.30(5)	541.95(28)	14.65(33)
	Chandra, 7 sources [23]	527.44(9)	541.72(18)	14.28(21)
	Chandra, shifted [23]	527.26(9)		
	Chandra, 11 sources [77]	527.39(2)		
	Chandra, 6 sources [78]	527.41(18)	541.77(40)	14.36(58)
Laboratory measurements	HZB [52]	527.26(4)	541.645(12)	
-	ALS [79,80]	526.79(4)	541.19(4)	14.40(8)
	ALS [76]	526.79(4)	541.20(4)	14.41(8)
	WSRC [81]	527.85(10)	541.27(15)	13.41(25)
	Auger spectroscopy [82]	527.20(30)		
MCHF converged result	MCHF [23]	527.49		

Table 1. Comparison of astronomically observed, laboratory measured, and computed O I K α and K β line energies (eV).

A further consideration in assessing the reliability of the ALS measurements was that the absolute energy calibration relied on earlier molecular measurements with uncertainty estimates that seemed questionable. Since the O I study by [23], new laboratory measurements have been performed using an electron beam ion trap (EBIT) to determine the K α resonance position at 527.26 \pm 0.04 eV [52], in good agreement with the observations and the converged MCHF result. This new measurement suggests a recalibration of the ALS reference spectrum.

4.3. Neon

For neon, the XSTAR atomic database includes the photoabsorption cross-sections by [13]. A benchmark for Ne II and Ne III was performed with *Chandra* and *XMM*-*Newton* observations of the bright low-mass X-ray binaries (LMXBs) Cygnus X–2 and XTE J1817-330 [35]. Column densities were obtained for the cold (Ne I) and warm (Ne II, Ne III) components of the ISM.

4.4. Magnesium

New photabsorption cross-sections for Mg species were computed by [25]. X-ray *XMM-Newton* spectra of the LMXB GS 1826-238 were used to perform a benchmark of the data. The resulting ionization fractions indicate that Mg is predominantly ionized rather than in neutral form, as also shown by UV observations.

4.5. Silicon

High-resolution *Chandra* spectra in the Si K-edge region (6–7 Å) from 16 LMXBs were analyzed by [39]. The Si model included the Si I photoabsorption cross-sections computed by Gorczyca et al. (2020, in preparation), while those for the ionized species were taken from [18]. The absorption features identified in the spectra agreed with the theoretical atomic data, even though the individual resonances of the K α "triplet" could not be resolved. Although a model without the dust component was used, good data fits were obtained, highlighting the need for accurate modeling of the gaseous component before attempting to address the solid component [39].

5. High-Density Effects on Atomic Parameters

XSTAR needs to address spectra from high-density sources such as the X-ray reflection from the inner region of the accretion disks around compact objects (electron temperature $T_e \sim 10^5 - 10^7$ K and density $n_e \sim 10^{18} - 10^{22}$ cm⁻³) [51,83]. This requires taking into account a series of atomic processes usually neglected in plasma photoionization models, which have been fully discussed by [46] and recently implemented in the code. We discuss in Sections 5.1 and 5.2 two of these processes—continuum lowering and DR suppression since they led to significant modifications of the database design and implementation.

5.1. Continuum Lowering

The density effects on the atomic thresholds (ionization potential and K edge), K α wavelengths and *A*-values, and Auger rates of oxygen and iron ions have been studied by [40–45]. The atom is assumed to be embedded in a weakly coupled plasma represented in the Multi-Configuration Dirac–Fock (MCDF) method with a time-independent Debye–Hückel screened Dirac–Coulomb Hamiltonian:

$$H_{DC}^{DH} = \sum_{i} c\vec{\alpha}_{i} \cdot \vec{p}_{i} + \beta_{i}c^{2} - \frac{Z}{r_{i}}e^{-\mu r_{i}} + \sum_{i>j} \frac{1}{r_{ij}}e^{-\mu r_{ij}}, \qquad (7)$$

where $r_{ij} = |\vec{r}_i - \vec{r}_j|$ and the plasma screening parameter μ (inverse of the Debye shielding length λ_D) is given in atomic units (λ_D in Bohr radius a_0 ; n_e in a_0^{-3} ; and kT_e in Hartree) by

$$\mu = \frac{1}{\lambda_D} = \sqrt{\frac{4\pi n_e}{kT_e}} \,. \tag{8}$$

As shown in Figure 8, the density effects mainly cause a threshold lowering that increases with μ , which for the ionization potential can be approximated in eV by the universal formula [45] in terms of the effective charge $Z_{\text{eff}} = Z - N + 1$

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$$\Delta E_0 = (-26.30 \pm 0.08) \mu Z_{\text{eff}} \,. \tag{9}$$



Figure 8. Ionization potential lowering in Fe IX–Fe XXV as a function of effective charge $Z_{\text{eff}} = Z - N + 1$ obtained with the MCDF method using a Debye–Hückel screening potential. Red circles: $\mu = 0.1$ a.u. Black circles: $\mu = 0.25$ a.u. Triangles: Debye–Hückel limit $\Delta E_0^{\text{DH}} = -27.2116 \,\mu Z_{\text{eff}}$ eV. Reproduced from Figure 1 of [43] with permission from Astronomy & Astrophysics, ©ESO.

In XSTAR continuum lowering leads to density-dependent targets with reduced numbers of levels, which must be derived at each density point causing a severe overhead at runtime. This bottleneck can be mitigated by pre-computing a μ grid of atomic databases but at the price of increasing the database volume significantly (~10 GB), thus hampering the standard package downloading procedure. Consequently, deployment strategies are being considered in terms of cloud virtual machines and containers.

5.2. DR Suppression

Rigorous collisional–radiative plasma modeling shows that DR is suppressed at high densities [84]. As mentioned in Section 2, RR and DR rates are treated separately in the XSTAR database using the AMDPP tabulation [64,65], which allows an estimate of the DR-rate density attenuation using the analytic formulae by [48,49]. The highly approximate value of the DR suppression factors has nevertheless been stressed by [49], as well as the need for detailed collisional–radiative calculations. The intermediate density range $(10 \le \log n_e \le 18 \text{ cm}^{-3})$ is expected to be the most susceptible since, at the higher densities $(\log n_e > 18 \text{ cm}^{-3})$, DR is practically negligible.

It is worth mentioning that in the unified treatment of electron–ion recombination [85,86], where the total recombination rate is obtained directly from a Maxwellian average of the photoionization cross-section through the Milne relation, it would be more difficult to evaluate the DR density effects. It would imply a rigorous approach to introduce plasma interactions in the Hamiltonian, such as those described for continuum lowering (see Section 5.1), followed by large-scale computations of photoionization cross-sections and fitting procedures to obtain a set of interpolation formulae.

To characterize the plasma effects caused by DR suppression, we have compared the heating and cooling rates in the ionization-parameter-temperature plane:

$$-3 \le \log(\xi) \le 3 \operatorname{erg\, cm s}^{-1}$$
 and $3.5 \le \log(T) \le 7.5 \operatorname{K}$ (10)

for XSTAR models with and without metal DR. We assume a power-law spectrum, density $n_e = 10^{10}$ cm⁻³, and luminosity $L = 10^{32}$ erg s⁻¹ (see Figure 9). The larger temperature differences occur in the interval $1 \le \log(\xi) \le 3$, where *T* is lower by as much as 40% when DR is excluded, and are caused by increases in the Fe average charge at $\log(\xi) \approx 1.3$ and

 $\log(\xi) \approx 2.5$. As shown in Figure 9 (right panel), the DR/noDR heating-rate ratio reaches a maximum at $(\log(\xi), \log(T)) \approx (1, 6.3)$, where the dominant species with and without DR are Fe XIII and Fe XVII, respectively. On the other hand, the cooling-rate ratio reaches a maximum at $(\log(\xi), \log(T)) \approx (-1, 5.4)$, caused by a higher oxygen average charge when DR is excluded.



Figure 9. Cooling- and heating-rate ratios with and without DR obtained from an XSTAR model with solar abundances, a power-law spectrum, density $n_e = 10^{10}$ cm⁻³, and luminosity $L = 10^{32}$ erg s⁻¹. Black dotted curve: thermal temperature with DR. Red dotted curve: thermal temperature without DR.

Concerning the Fe anomalous overabundance derived from the reflection spectra of accretion disks [51], we have computed the X-ray spectra reflected from an optically thick atmosphere using the XILLVER code [87,88]. The atmosphere is assumed to have plane-parallel geometry and a constant gas density of $\log(n_e) = 18 \text{ cm}^{-3}$. The ionization parameter is set to $\xi = 4\pi F_x/n_e = 32 \text{ erg cm} \text{ s}^{-1}$ implying an illuminating flux of $F_x = 3 \times 10^{18} \text{ erg cm}^{-2} \text{ s}^{-1}$. The ionization balance is calculated self-consistently using the routines and atomic data from XSTAR, setting all elemental abundances to their solar values. It is found that metal DR suppression leads to significant changes in the ionization state of the gas resulting in more ionized species. As shown in Figure 10, there is a strong enhancement of the Fe XVII L lines and Fe XVIII K lines.



Figure 10. Reflected X-ray spectrum from an optically thick atmosphere modeled using XILLVER with and without DR. A plane-parallel atmosphere with solar abundances and density $\log(n_e) = 18 \text{ cm}^{-3}$ was assumed.

It is worth mentioning that continuum lowering is not taken into account in this XILLVER test calculation since it is expected to have a noticeable impact only at densities $n_e > 10^{19}$ cm⁻³. The calculation was performed at $n_e = 10^{18}$ cm⁻³ to show that DR effects become conspicuous at lower densities. Further characterization of high-density effects are reported in [46].

Density effects on DR and the high-*n* levels through continuum lowering are taken into account in the XSTAR database via the rates into one or two superlevels for each ion. The recombination rate onto the superlevel is tabulated on a density–temperature grid and stored in data types 70 and 99 (see Appendix A). For the former, rates are tabulated on a log scale, while for the latter, they are tabulated directly. The total recombination rate onto the superlevels plus the rates onto the other levels that are treated explicitly; these are called the "spectroscopic levels". The recombination rates onto the superlevels are chosen such that, when they are used to calculate a recombination rate and all the rates are summed up, the total rate taken from one of the AMDPP compilations [64] where available or from [89] otherwise. We apply density-dependent suppression factors to the recombination rates onto the superlevel to take into account density effects on radiative and dielectronic recombination. They are derived from a general formula that depends on the isoelectronic sequence, charge, density, and temperature [48,49].

The superlevels decay directly to the ground level without the emission of any observable cascade radiation for ions with three or more electrons. The exception is the decay of members of the H and He isoelectronic sequences, for which we use explicitly calculated cascade matrix calculations to treat the decay of the superlevels to the spectroscopic levels [90,91]. The superlevels are chosen to have energies close to the continuum. We include both radiative and collisional transitions to the ground level (and other levels in the case of H- and He-like ions).

6. Database Curation

The recent inclusion in the XSTAR database of atomic parameters for odd-*Z* elements, Z = (9, 11, 15, 17, 19), and trace elements with Z = (21-25, 27, 29, 30) [26–29] practically doubled the database volume when compared with previous versions that used hydrogenic scaling for these elements, bringing to the fore a series of curatorial problems. The database upgrade was thus lengthy and demanding in detail. In Sections 6.1–6.4, we give a brief account of some the issues encountered and steps taken to solve them.

6.1. Atomic Models

As shown in Table 9 of [26], the target models for ions with atomic numbers Z = (21-25, 27, 29, 30) and electron number $19 \le N \le Z - 2$ include levels from the valence configurations $\mu = 3d^x 4s^y$ and the L- and K-vacancy configurations $3p^{-1}\mu^{+1}$, $2p^{-1}\mu^{+1}$, and $1s^{-1}\mu^{+1}$. The total number of target levels in each of these species is larger than 500 and in some cases (e.g., N = 23) close to 2000. It was found that to model astrophysical plasmas in XSTAR with such large atomic representations was impractical and with little tangible gains. We thus excluded the levels belonging to μ configurations containing 4s orbitals—specifically, y = 0 in μ for N > 19. This decision led to a reduction in the number of target levels to a few hundreds and implied extensive level renumbering for these sequences.

6.2. Metastable Levels

As discussed in Section 2, the XSTAR atomic models display occasional low-lying metastable levels (e.g., the 3s1.3p1.3P_0 level in Mg-like ions) with no radiative decay transitions. Since metastable lavels in low valence-shell configurations are astrophysically important, alternative decay mechanisms such as collisional de-excitation must be specified for such levels. For instance, we computed a large number of distorted-wave collision strengths with AUTOSTRUCTURE for transitions within the valence-shell configurations of trace-element ions (see record with ID = mb01 and code = 1150 in Table A2). However, for ions with electron number N > 19 and ground configuration $3d^x$, atomic models with a few hundred levels must be considered (see Section 6.1), which leads to the appearance of unexpected metastable levels in the $3p^{-1}$ configuration at fairly high energies. Most of these levels were detected by running the code to look for levels with anomalously large populations, which were then removed by hand followed by level reordering.

6.3. Database Pointers

As previously mentioned, the raw XSTAR database consists of a list of ASCII files labeled diont0dt, where ion is the ion identifier and dt the data type. When the package is released, these files are read into three one-dimensional arrays (integer, real, and character variables) and a set of pointers to be finally structured into the FITS file atdb.fits. At runtime, these arrays are read into main memory to be accessed according to the user model. This data structure arrangement is efficient when uploading from disk and during processing, but can be cumbersome in database updating when new record types are created. Proficient nursing and testing are required before the new version is released.

6.4. Metadata

The XSTAR database compiles atomic data computed to order and from a myriad of other data sources. The data provenance of the first release was detailed in [2], but since then, the database has undergone several major upgrades that have not been formally registered within its file structure. One of the aims of the present paper is to create an up-to-date data provenance blueprint to be included in the database using the FITS metadata facilities.

In Table A2 of Appendix C, we have tabulated an up-to-date list of data source references including the original four- and five-letter mnemonics used for identification in [2] and a new four-integer code to be inserted in every database record. For this purpose, we will adopt the third integer (currently displaying a zero) of the record header $(dt, rt, 0, N_{\text{flt}}, N_{\text{int}}, N_{\text{chr}})$ described in Section 2 to denote this data provenance code.

6.5. Relational Integrity

As described in Section 2, the data in each diont0dt file are self-contained to simplify and speed up the calculation of a wide variety of rates at runtime, but as a result, this scheme obviates the relational integrity of the database. For instance, the wavelengths displayed in file diont050 (see Figure 3) are not automatically recalculated if level energies are updated in file diont006 (Figure 2). The level attributes specified in the latter file are also repeated in several other data types, e.g., the K-vacancy level energy in file diont086 (Figure 4). The extent of this complication is further illustrated in the list of data types in Appendix A. Data type 14 lists the ionization potential for every ionic species, but this datum is also contained in the floating-point tuple of every level in data type 06. Some of the level integer attributes of the latter data type are included in data types dealing with the effective ion charge (57), photoionization cross-sections (49, 53, 85, and 88), superlevel recombination rates (70 and 99), and satellite-level autoionization rates (72).

This database model can therefore make updating an involved and error-prone process, which can be mitigated by adopting an SQL-based relational database engine and devising a utility to perform the transcription to the atdb.fits file. Despite the wide variety of data types (see Appendix A) and record lengths, this is possible as shown by the development of the uaDB database (see Section 8). However, this scheme would hamper the generation of new data types, which is central to the XSTAR data curation strategy (see Section 2), and also lead to a more voluminous atdb.fits file. This proposition is being considered in the XSTAR development roadmap.

6.6. Atomic Data Accuracy

As astronomical observations become ever more accurate (see, for example, Figure 1), the reliability of the atomic data used in spectral modeling is under recurrent scrutiny, and hence, data assessment activities are key. In this respect, we have seen three schemes: by a reputed institution such as NIST [92]; by appointed review panels (e.g., VAMDC [93]), and by an open community of both data producers and users (e.g., AtomPy [94]). Our research group comprises both computational atomic physicists and astrophysical modelers, so we favor the latter. However, it has not been easy to determine accuracy ratings in our datasets for K-line diagnostics (Section 3) due to the lack of laboratory measurements and

despite adopting a multi-code computational strategy. As described in Section 4, we have pursued benchmarks of the photoabsorption cross-sections using astronomical observations of ISM K absorption, but they led to a polemic between laboratory spectroscopists and observational astronomers that was fortunately settled.

We have mentioned in Section 2 that our main concern regarding the XSTAR database upgrade has been the complete datasets for K-shell processes in ions with $Z \leq 30$, at the expense of neglecting those for the valence-shell that in general imply data collection rather than new computations. It may be seen in Appendix C that most of the current datasets for valence-shell transitions go back to the original version [2] then demanding an extensive overhaul. This is one of our main priorities in the database curation.

7. ISMabs

Although the main components of the ISM are neutral ionic species, the also perceived existence of charged ions prompted us to develop a new ISM X-ray absorption model referred to as ISMabs [35]. It includes atomic data for singly and doubly ionized species of cosmically abundant elements, namely H, He, C, N, O, Ne, Mg, Si, S, Ar, Ca, and Fe, in addition to the neutral systems. This model allows ion column densities to be determined directly, and the inclusion of ionized species has led to improved spectral fits as compared to those only considering neutral systems. The completeness and accuracy of the relevant atomic data are crucial to avoid misidentification and misinterpretation of the absorption features detected in analyses of ISM X-ray absorption spectra relying on fits of Gaussian profiles. We have performed with this model a detailed study of X-ray absorption in the local ISM by analyzing spectra from 24 Galactic sources. We have estimated the fractions of neutral and singly and doubly ionized species of O, Ne, and Fe, confirming the dominance of the cold component and a pervasive low degree of ionization [36].

8. Universal Atomic Database (uaDB)

The collection of atomic data in XSTAR is comprised of data in many different formats from many different sources. To assist in disseminating these data to the larger community and to account for the addition of new data, the Universal Atomic Database (uaDB) was developed. The uaDB is a mySQL database hosted at NASA/GSFC, accessed via its website (https://heasarc.gsfc.nasa.gov/uadb/index.php), and data can be queried based on a number of properties such as ion and data type. The web page also contains tools to help assess what data are available across ions or data types. One such tool displays the type of data available on a grid of element vs. ion (see Figure 11) while another displays all atomic lines present in a given wavelength range.

The main features of uaDB are:

- Each dataset contains a reference to the source paper(s);
- Data are stored in the original format and units as published;
- Multiple entries can exist for the same quantity (e.g., theoretical vs. measured energy levels);
- The coupling scheme of atomic states is built into the database structure;
- Data collections exist that define a complete model; no duplicate data exist within a collection.

All data in uaDB have a reference to the data source. The reference consists of a link to the journal article where the data were obtained and to its entry in NASA/ADS. The user is never more than a couple mouse clicks away from reading the source article. Data are entered into uaDB in the same format and with the same units as published. This means that, for example, some energy levels are stored in eV and others in cm⁻¹. The uaDB understands these units and can internally convert them for data querying in an energy range. The uaDB can also convert between quantities that differ by a statistical weight (e.g., *f*-values and *gf*-values) since it understands the level structure. We do not modify published data in order to be as true as possible to the original source and to help identify transcription



errors. However, this can make it more difficult to compare data from different sources. To help compare data, tools are developed that can perform the necessary conversions.

HEASARC Home | Observatories | Archive | Calibration | Software | Tools | Students/Teachers/Public

Figure 11. Grid tool in uaDB.

When compiling a set of atomic data for modeling purposes, it is important to have coverage; that is, for each state included in the model, a full set of data connecting that state with other states is necessary. If there are data allowing the population of a state to increase but no data to decrease it, then the model as a whole will be flawed. Furthermore, if the atomic model encompasses a large set of ions, it is difficult to obtain atomic data from the literature of equal quality. We often need to combine level-resolved data with term-resolved data to achieve coverage for an ion. To help with these difficulties, uaDB has different coupling schemes built into its structure and is aware of how different coupling schemes relate. Thus, when combining *LS* states with *LSJ*, uaDB can detect if any states are being double-counted. There are also routines for converting data from one coupling scheme to another based on simple statistical weighting.

The uaDB currently contains the complete set of data included in XSTAR as well as the level data from CHIANTI. To assist in distinguishing these data, uaDB defines collections. A collection is a dataset where there is no duplication of values for the same quantity. Currently, in uaDB the collection is just a label that can be used to constrain a query, but in the future collections could be created dynamically to construct a custom model. Collections are not part of the XSTAR development program, but have been widely used in, for instance, the PyNeb Python package to model emission lines in gaseous nebulae and, more recently, to assess atomic data accuracy [95,96].

9. Discussion and Conclusions

The present report describes in detail the updated version of the underlying atomic database of the XSTAR photoionization code to replace the original implementation of [2]. We include complete lists of the current rate types, data types (Appendix A), ionic models (Appendix B), and source references (Appendix C). We summarize the systematic calcula-

tions performed to characterize the metal K-lines for astronomical X-ray spectral modeling and recent extensions of the database, namely continuum lowering and dielectronic recombination suppression, to deal with high-density ($n_e \leq 10^{22}$ cm⁻³) plasmas. We have tried to convey the intricacies in the development and maintenance of an application-based atomic database, which go way beyond the compilation of the data and lead to curatorial and integrity problems caused by the ever-increasing dataset volumes.

An important finding is that the atomic models used to compute the data are not necessarily suitable for plasma modeling, particularly for ionic species with electron number N > 19. Level-number trimming was necessary to implement functional atomic representations for such systems, although their soundness in non-LTE modeling deserves further attention as the appearance of unexpected metastable levels becomes an issue.

A further point worth emphasizing is that the XSTAR atomic database and output files are structured under the FITS format specifications, which has been established as the standard for astronomical datasets associated to the new generation of telescopes. We believe that this compliance will reduce user stress in intense data processing, which is bound to take place at the data repositories rather than locally.

We have briefly recounted two projects that have emerged from the XSTAR database, namely the ISMabs absorption model and the uaDB database. The former has become an important initiative as it has led to benchmarks of the atomic data with observations and laboratory measurements; however, it has not been a smooth process as unexpected inaccuracies in the calibration of the laboratory wavelength scale were detected and finally resolved [23,52]. This outcome has given us some confidence in the identification of the spectral features and the accuracy of our atomic data. The uaDB database model will be helpful in future projects to disseminate the larger atomic data volumes required in non-LTE modeling.

A question that remains unanswered is whether our efforts to ensure accuracy and completeness for the XSTAR database are sufficient to exploit the high resolution and sensitivity expected from the microcalorimeter-based spectrometers aboard the new X-ray space telescopes. The encouraging fit of the showcase HITOMI spectrum of Figure 1 required basic data manipulation, but at least it was conclusive. Therefore, we envision further database refinement in the near future, especially improved datasets for valence-shell processes, to provide a reliable modeling platform for high-resolution spectroscopy in the new era. In this respect more laboratory benchmarks would be welcome.

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Data Availability Statement: The XSTAR modeling code, the uaDB database, and the ISMabs model are available from the website of the NASA High Energy Astrophysics Science Archive Research Center (HEASARC, https://heasarc.gsfc.nasa.gov/).

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Abbreviations

The following abbreviations are used in this manuscript:

ADAS	Atomic Data and Analysis Structure
ADS	Astrophysics Data System
ALS	Advanced Light Source
AMDPP	Atomic and Molecular Diagnostic Processes in Plasmas
FITS	Flexible Image Transport System
GSFC	Goddard Space Flight Center
HZB	Helmholtz-Zentrum Berlin
IPNAS	Institut de Physique Nucléaire, Atomique et de Spectrométrie
LTE	Local Thermodynamic Equilibrium
NASA	National Aeronautics and Space Administration
NIST	National Institute of Standards and Technology
PP95	Atomic Data for the Analysis of Emission Lines
SQL	Structured Query Language
WSRC	Wisconsin Synchrotron Radiation Center

Appendix A. Data Types

This appendix lists the data types of the XSTAR database. Unless otherwise stated, the temperature *T* is given in 10^4 K units. The record attributes are: r1, r2, ... for the real variables; i1, i2, ... for the integer variables; and s1, s2, ... for the character strings. The following notation is used: *ion* is the ion identifier; *Z* the atomic number; *N* the electron number; $E(\infty)$ the ionization potential; E(i) the *i*th level energy; *n* the principal quantum number; ν the effective quantum number; *S* the total spin quantum number; *L* the total orbital angular momentum quantum number; *J* the total angular momentum quantum number; *a* the radiative transition wavelength; gf(i, k) the weighted oscillator strength; and A(k, i) the transition rate.

- **01.** Radiative recombination rate coefficient of *N*-electron recombined ion [89,97]: $r1 = A_{rad}$ (cm³ s⁻¹); $r2 = \eta$; $i1 = ion_N$.
- **02.** H⁰ charge exchange rate coefficient of *N*-electron recombined ion [98]: r1 = a $(10^{-9} \text{ cm}^3 \text{ s}^{-1})$; r2 = b; r3 = c; r4 = d; $r5 = T_1$ (K); $r6 = T_2$ (K); $r7 = \Delta E/k$ (10^4 K) ; $i1 = ion_N$; s1 = recombining ion identifier.
- **06.** Data attributes of the *i*th level of *N*-electron ion: r1 = E(i) (eV); r2 = (2J + 1); $r3 = \nu$ (effective quantum number); $r4 = E(\infty)$ (eV); i1 = n; i2 = (2S + 1); i3 = L; i4 = Z; i5 = i; $i6 = ion_N$; s1 = level assignment (electron configuration and spectroscopic term).
- **07.** Dielectronic recombination rate coefficient of *N*-electron recombined ion [89,97]: $r1 = A_{di}$ (cm³ s⁻¹ K^{3/2}); $r2 = B_{di}$; $r3 = T_0$; $r4 = T_1$; $i1 = ion_N$.
- 14. Ionization potential of *N*-electron ion: $r1 = E(\infty)$ (eV); i1 = Z N + 1; i2 = Z; $i3 = ion_N$; s1 = ion identifier.
- **22.** Dielectronic recombination rate coefficient of the *N*-electron recombined ion [99]: r1 = a; r2 = b; r3 = c; r4 = d; r5 = f; $i1 = ion_N$.
- **30.** Total radiative recombination rate (hydrogenic) for *N*-electron recombined ion [100]: i1 = Z; $i2 = ion_N$.
- **38.** Total radiative recombination rate coefficient of *N*-electron recombined ion [http: //amdpp.phys.strath.ac.uk/tamoc/DATA/RR/]: $r1 = A (cm^3 s^{-1})$; r2 = B; $r3 = T_0$ (K); $r4 = T_1$ (K); r5 = C; $r6 = T_2$ (K); i1 = Z; i2 = N - 1; i3 = M; i4 = W; $i5 = ion_N$.

- 39. Total dielectronic recombination rate coefficient of N-electron recombined ion [http://amdpp.phys.strath.ac.uk/tamoc/DATA/DR/]: r1-rjmax = (C(j), j = 1, jmax) (cm³ s⁻¹ K^{3/2}); rjmax+1-rj_{2*max} = (T(j), j = 1, jmax) (K); i1 = Z; i2 = N 1; i3 = M; i4 = W, i5 = ion_N.
- **49.** Partial photoionization cross-section of i_N th level of the *N*-electron ion leaving the (N-1)-electron ion in the k_{N-1} th level: $r1-rj_{2*max} = (E(j), \sigma(E(j)), j = 1, j_{max})$ (Energy in Ryd relative to $E(\infty)$, cross-section in Mb); i1 = n; i2 = L; i3 = 2J; i4 = Z; $i5 = k_{N-1}$; $i6 = ion_{N-1}$; $i7 = i_N$; $i8 = ion_N$.
- **50.** Line (k i) radiation rates of *N*-electron ion: $r1 = \lambda$ (Å); r2 = gf(i,k); r3 = A(k,i) (s⁻¹); i1 = i (lower level); i2 = k (upper level); i3 = Z; $i4 = ion_N$.
- **51.** Electron-impact effective collision strength for the k i transition of *N*-electron ion (CHIANTI fit [61,101]): r1 = ΔE (Ryd); r2 = C; r3-r7 = (Y_{red}(*j*), *j* = 1,5) (reduced effective collision strength); i1 = *it* (transition type); i2 = *i* (lower level); i3 = *k* (upper level); i4 = *Z*; i5 = *ion*_N.
- **53.** TOPbase partial photoionization cross-section (resonance averaged) of i_N th level of the *N*-electron ion leaving the (N 1)-electron ion in the k_{N-1} th level: $r1 rj_{2*max} = (E(j), \sigma(E(j)), j = 1, j_{max})$ (Energy in Ryd relative to $E(\infty)$, cross-section in Mb); $i1 = n; i2 = L; i3 = 2J; i4 = Z; i5 = k_{N-1}; i6 = ion_{N-1}; i7 = i_N; i8 = ion_N.$
- 54. Radiative transition probability A_{ki} for the k i transition of *N*-electron ion computed by quantum defect theory (or hydrogenic): r1 = 0.0E+0; i1 = i (lower level); i2 = k (upper level); i3 = Z; $i4 = ion_N$.
- **56.** Electron-impact effective collision strengths for the k i transition of *N*-electron ion: $r1-rj_{max} = (\log T_e(j), j = 1, j_{max})$ (K); $rj_{(max+1)} - rj_{(2*max)} = (Y(T_e(j)), j = 1, j_{max})$ (effective collision strength); i1 = i (lower level); i2 = k (upper level); i3 = Z; $i4 = ion_N$.
- **57.** Effective ion charge for *i*th level of *N*-electron ion used in collisional ionization rates: $r1 = Z_{eff}$; i1 = n; i2 = L; i3 = 2J; i4 = i; $i5 = ion_N$
- **59.** Partial photoionization cross-section of i_N th level of the *N*-electron ion leaving the (N-1)-electron ion in the k_{N-1} th level [102]: r1 = E(th) (eV); r2 = E(0) (eV); $r3 = \sigma(0)$ (Mb); r4 = y(a); r5 = P; r6 = y(w); i1 = N; i2 = n (shell principal quantum number); i3 = l (subshell orbital quantum number); $i4 = k_{N-1}$; $i5 = ion_{N-1}$; $i6 = i_N$; $i7 = ion_N$; s1 = shell-ion identifier.
- 60. Analytic fits for effective collision strengths in H-like ions [103]: r1-rj_{max} = coefficients; i1 = *i* (lower level); i2 = *k* (upper level); i3 = *nphot*; i4 = *ion*_N; s1 = Transition.
- 62. Analytic fits for effective collision strengths in H-like ions [103]: r1-rj_{max} = coefficients; i1 = *i* (lower level); i2 = *k* (upper level); i3 = 1; i4 = *i*on_N; s1 = Transition.
- **63.** Collisional transition probability C_{ik} for *N*-electron ion computed by quantum defect theory (or hydrogenic): i1 = 1 i2 = i (lower level); i3 = k (upper level); i4 = Z; $i5 = ion_N$.
- 66. Fits to fine-structure collision strengths for He-like ions [104]: r1-rj_{max} = coefficients;
 i1 = *i* (lower level); i2 = *k* (upper level); i3 = *Z*; i4 = *ion*_N.
- 67. Analytic fits for effective collision strengths in He-like ions [105]: r1-rj_{max} = coefficients; i1 = *i* (lower level); i2 = *k* (upper level); i3 = *Z*; i4 = *ion*_N.
- 68. Analytic fits for effective collision strengths in He-like ions [106]: r1-rj_{max} = coefficients; i1 = *i* (lower level); i2 = *k* (upper level); i3 = *Z*; i4 = *ion*_N.
- 69. Fits to LS collision strengths for He-like ions [104]: r1-rj_{max} = coefficients; i1 = i (lower level); i2 = k (upper level); i3 = Z; i4 = ion_N.
- 70. Coefficients for recombination and photoionization cross-sections of superlevels: $r1-rj_{nd} = (n_e(j), j = 1, j_{nd}) (cm^{-3}); rj_{nd+1}-rj_{nd+nt} = (T_e(j), j = 1, j_{nt}); rj_{nd+nt+1} - rj_{nd+nt+nt*nd} = ((\log \alpha(j, j'), j = 1, j_{nt}), j' = 1, j'_{nd}); rj_{nd+nt+nt*nd+1} - rj_{nd+nt+nt*nd+2*nx} = (E(j), \sigma(j), j = 1, j_{nx}) (Ryd, Mb); i1 = nd; i2 = nt; i3 = nx;$
- i4 = n; i5 = L; i6 = 2S + 1; i7 = Z; i8 = k_{N-1} ; i9 = ion_{N-1} ; i10 = i_N ; i11 = ion_N . 71. Radiative transition rates from superlevels to spectroscopic levels: r1-rj_{nd} = $(n_e(j), j = 1, j_{nd})$ (cm⁻³); rj_{nd+1}-rj_{nd+nt} = $(T_e(j), j = 1, j_{nt})$; rj_{nd+nt+1}-rj_{nd+nt+nt*nd} =

 $((A(j,j'), j = 1, j_{nt}), j' = 1, j'_{nd})$ (s⁻¹); $r_{nd+nt+nt*nd+1} = \lambda$ (Å); i1 = nd; i2 = nt; i3 = *i* (lower level); i4 = *k* (upper level); i5 = *Z*; i6 = *ion*_N.

- 72. Autoionization rates for satellite levels: $r1 = A_a(k,i)$ (s⁻¹); r2 = E(k) (eV above ionization limit); r3 = (2J + 1); i1 = (2S + 1); i2 = L; i3 = k (level); i4 = i (continuum level); i5 = Z; $i6 = ion_N$; s1 = level configuration.
- **73.** Fit to effective collision strengths for satellite levels of He-like ions [107]: r1-r7 = fit coefficients; i1 = i (lower level); i2 = j (upper level); i3 = Z; $i4 = ion_N$.
- 74. Delta functions to add to photoionization cross-sections to match ADAS DR rates: $r1 = E(\infty)$ (Ryd); $r2-rj_{m+1} = (E(j), j = 1, j_m)$ (Ryd); $rj_{m+2}-rj_{2*m+1} = (f(j), j = 1, j_m)$ (cm²); i1 = n; i2 = L; i3 = 2S + 1; i4 = Z; $i5 = k_{N-1}$; $i6 = ion_{N-1}$; $i7 = i_N$; $i8 = ion_N$.
- **75.** Autoionization rates for Fe XXIV satellites [5]: $r1 = A_a(k,i)$ (s⁻¹); r2 = E(k) (eV above ionization limit); $i1 = ion_N$, $i2 = k_N$; $i3 = ion_{N-1}$; $i4 = i_{N-1}$; $i5 = ion_N$.
- **76.** Two-photon radiation rate for (k i) transition of *N*-electron ion: r1 = A(k, i) (s⁻¹); i1 = *i* (lower level); i2 = *k* (upper level); i3 = *N*; i4 = *ion*_N; s1 = transition.
- 77. Collision transition rates from superlevels to spectroscopic levels: $r1-rj_{nd} = (n_e(j), j = 1, j_{nd})$ (cm⁻³); $rj_{nd+1}-rj_{nd+nt} = (T_e(j), j = 1, j_{nt})$; $rj_{nd+nt+1}-rj_{nd+nt+nt*nd} = ((C(j, j'), j = 1, j_{nt}), j' = 1, j'_{nd})$ (s⁻¹); $rj_{nd+nt+nt*nd+1} = \lambda$ (Å); i1 = nd; i2 = nt; i3 = i (lower level); i4 = k (upper level); i5 = Z; $i6 = ion_N$.
- **81.** Collision strengths for Fe XIX [108]: r1 = Y(k, i); i1 = i (lower level); i2 = k (upper level); i3 = Z; $i4 = ion_N$.
- **82.** Decay rates for Fe UTA [109]: $r1 = \lambda$ (Å); r2 = E(k) (eV); r3 = gf(i,k); $r4 = A_r(k,i)$ (s⁻¹); $r5 = A_a(k,i)$ (s⁻¹); i1 = i (lower level); i2 = k (upper level); $i3 = ion_N$.
- **83.** Level data for Fe UTA [109]: r1 = E(i) (eV); r2 = (2J + 1); r3 = 0.0; r4 = 0.0; i1 = 1; i2 = i (level); $i3 = ion_N$; s1 = level configuration assignment.
- **85.** Photoionization cross-sections for Fe ions obtained by summation of resonances near the K edge [3]: $r1 = Z_{eff}$; $r2 = E_{th}$ (Ryd); r3 = f; $r4 = \gamma$; r5 = scaling factor; i1 = n; i2 = L; i3 = 2J; i4 = Z; $i5 = k_{N-1}$; $i6 = ion_{N-1}$; $i7 = i_N$; $i8 = ion_N$.
- **86.** Auger and radiative widths of k_N th K-vacancy level: $r1 = E(k_N)$ (eV, relative to $E(\infty)$); $r2 = A_a(k_N)$ (s⁻¹); $r3 = A_a(k_N, i_{N-1})$ (s⁻¹); $r4 = A_r(k_N)$ (s⁻¹); $i1 = i_{N-1}$; $i2 = k_N$; i3 = Z; $i4 = ion_{N-1}$; $i5 = ion_N$.
- **88.** Photoionization cross-section damped excess of i_N th level of the *N*-electron ion leaving the (N 1)-electron ion in superlevel_[K] k_{N-1} : r1-rj_{2*max} = $(E(j), \sigma(E(j)), j = 1, j_{max})$ (Energy in Ryd relative to $E(\infty)$, cross-section in Mb); i1 = n; i2 = L; i3 = 2J; i4 = Z; i5 = k_{N-1} ; i6 = i_N ; i7 = ion_N .
- **91.** APED line (k i) radiation rates [110]: $r1 = \lambda$ (Å); r2 = 0.0; r3 = A(k, i) (s⁻¹); i1 = i (lower level); i2 = k (upper level); i3 = Z; $i4 = ion_N$.
- **92.** APED collision strengths [110]: $r1 = T_e(1)$ (K); $r2 = T_e(j_{max})$ (K); $r3-rj_{max+2} = (T_e(j), j = 1, j_{max})$ (K); $rj_{max+3}-rj_{2*max+2} = (Y(j), j = 1, j_{max})$; i1 = i (lower level); i2 = k (upper level); i3 = 113; i4 = Z; $i5 = ion_N$.
- **95.** Collisional ionization rates for *N*-electron ion [111]: r1 = E(th) (eV); $r2 = T_0$ (K); $r3-rj_{max+2} = (\rho(j), j = 1, j_{max})$ (effective collision strength); i1 = i (level); $i5 = ion_N$.
- **98.** Electron-impact effective collision strengths for the k i transition of the *N*-electron ion (CHIANTI fit [61,101]): r1 = ΔE (Ryd); r2 = C; r3-rj_{max+2} = (Y_{red}(*j*), $j = 1, j_{max}$) (reduced effective collision strength); i1 = *i*; i2 = *k*; i3 = *it* (transition type); i4 = *ion*_N.
- **99.** Coefficients for recombination and photoionization cross-sections of superlevels: $r1-rj_{nd} = (n_e(j), j = 1, j_{nd}) (cm^{-3}); rj_{nd+1}-rj_{nd+nt} = (T_e(j), j = 1, j_{nt}); rj_{nd+nt+1} - rj_{nd+nt+nt*nd} = ((\alpha(j, j'), j = 1, j_{nt}), j' = 1, j'_{nd}); rj_{nd+nt+nt*nd+1} - rj_{nd+nt+nt*nd+2*nx} = (E(j), \sigma(j), j = 1, j_{nx}) (Ryd, Mb); i1 = nd; i2 = nt; i3 = nx; i4 = n; i5 = L; i6 = 2S + 1; i7 = Z; i8 = k_{N-1}; i9 = ion_{N-1}; i10 = i_N; i11 = ion_N.$

Appendix B. Ion Models

Z N	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15
1	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33
2		46	32	32	32	56	58	241	52	54	52	58	52	58	52
3			3	3	3	26	43	43	22	43	22	43	22	43	22
4				3	3	22	53	53	43	79	43	79	43	79	43
5					3	22	58	163	53	163	53	163	53	163	53
6						8	61	79	53	79	53	79	53	79	53
7							82	34	34	34	34	34	63	34	34
8								19	3	19	19	19	57	19	19
9									3	118	7	7	7	7	7
10										4	3	46	46	46	50
11											3	24	4	60	42
12												3	4	57	31
13													3	18	55
14														3	3
15															3
N Z	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30
1	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33
2	58	52	38	52	46	52	52	52	52	52	50	52	43	52	52
3	43	22	43	22	43	22	22	22	22	22	52	22	53	22	22
4	79	43	79	43	79	43	43	43	43	43	199	43	79	43	43
5	163	53	163	53	163	53	53	53	53	53	551	53	163	53	53
6	79	53	53	53	53	53	53	53	53	53	623	53	53	53	53
7	34	34	34	34	34	34	34	34	34	34	744	34	34	34	34
8	19	19	19	19	29	19	19	19	19	19	639	19	67	19	19
9	7	7	7	7	7	7	7	7	7	7	340	7	117	7	7
10	46	50	72	50	78	50	50	50	50	50	276	50	98	50	50
11	60	42	208	42	208	42	42	42	42	42	85	42	216	42	42
12	53	31	153	31	149	31	31	31	31	31	146	31	1/1	31	31
13	/0	55 71	208	55 71	208	55 71	55 71	55 71	55 71	55 71	55	55 71	235	55 71	55 71
14	82 21	/1	249	/1	249	/1	71	71	71	71	68 79	71	269	/1 EE	71
15	21	55	211 142	21	211 142	21	21	21	21	21	70	21	219 145	21	21
10	3	3	145 6 2	51 10	145 62	51 10	51 10	51 10	51 10	51 10	62	51 10	145 62	51 10	51 10
17		3	02 2	10	02 2	10 54	10 54	10 54	10 54	10 54	02 21	10 54	40	10 54	10 54
10			3	3	5	221	221	221	221	221	170	221	40 1/3	221	24
20				5	3	32	258	258	258	258	54	258	258	258	221
20					5	3	200	441	230	441	55	441	441	441	230 441
21						5	3	3	537	537	33	537	537	537	537
22							5	3	163	463	33 70	463	463	463	<i>1</i> 63
23								5	3	160	73	295	295	295	295
25									5	3	197	128	128	128	128
25										5	3	3	40	40	40
20											5	3	т0 2	<u>२</u>	т 0 २
28												5	3	3	3
20													5	3	3
30														0	3
															5

Appendix C. Atomic Data Provenance

Table A2. Data provenance including the CHIANTI [61], PP95 [63], and AMDPP [64,65] compilations. rt = rate type. dt = data type.

ID	Code	rt	dt	(Z; N)	References
ac81	1000	3	56	(8;1)	[112]
ac92	1001	3	56	(2;1)	[113]
ak91	1002	3	56	(6;1)	[114]
ak91a	1003	3	56	(10; 1)	[115]
ak92	1004	3	56	(14; 1)	[116]
ak92a	1005	3	56	(20; 1)	[117]
ak93	1006	3	56	(26; 1)	[118]
ba00	1007	4	50	(28;2)	[91]
bm98	1008	3	56	(26;15)	[119], CHIANTI ([120,121])
bn00	1009	3	56	(26; 20)	[122], CHIANTI ([123])
bp98	1010	3	56	(26; 25)	[90]
ch00	1011	3	51	(10;7), (20;8), (26;16), (28;14)	CHIANTI
ch00	1012	4	50	(6,7,8;4), (12,14;5), (20;8),	CHIANTI
				(26; 6, 12, 14-18), (28; 14)	
ch01	1013	3	51	(6.7:3)	CHIANTI ([124])
ch01	1014	4	50	(6,7;3)	CHIANTI ([125,126])
ch02	1015	4	50	(8, 10, 12, 14, 16, 18, 20, 26, 28; 3)	CHIANTI ([126,127])
ch02	1016	3	51	(8, 10, 12, 14, 16, 18, 20, 28; 3)	CHIANTI ([127])
ch04	1017	3	51	(6:4)	CHIANTI ([128,129])
ch04	1018	4	50	(10, 12, 16; 4)	CHIANTI ([130–132])
ch05	1019	3	51	(7:4)	CHIANTI ([133])
ch05	1020	4	50	(14, 18, 26; 4)	CHIANTI, CHIANTI ([131])
ch06	1021	3	51	(8:4)	CHIANTI $([132, 134])$
ch06	1022	4	50	(20, 28; 4)	([130,132])
ch07	1023	3	51	(10, 12, 14, 16, 18, 20, 26, 28; 4)	([131,132])
ch07	1024	4	50	(6:5)	CHIANTI ($[135-137]$) NIST
ch08	1025	3	51	(6;5)	CHIANTI ([138])
ch08	1026	4	50	(7.5)	(139 140]
ch09	1027	4	50	(8, 10; 5)	CHIANTI ([135])
ch09	1022	3	51	(7.5)	CHIANTI ([141 142])
ch10	1020	4	50	(16, 18, 20; 5)	([11], [12])
ch10	1020	3	51	(10, 10, 20, 0) (8 10 12 14 16 18 20 26:5)	CHIANTI ([144])
ch11	1031	3	51	(28:5)	CHIANTI ([145])
ch11	1032	4	50	(26;5)	CHIANTI ([145])
ch12	1032	- - -	50	(28:5)	([135] 143])
ch12	1034	3	51	(7:6)	([130,140,140])
ch13	1035	4	50	(7,6)	CHIANTI ([141,142])
ch13	1036	3	51	(8:6)	(110)
ch14	1037	4	50	(8,6)	CHIANTI ([149])
ch14	1038	3	51	(10:6)	(1150-152)
ch15	1039	4	50	(10,0) (10,12,14.6)	$([150 \ 152])$
ch15	1037	2	51	(12:6)	CHIANTI ([152,155])
ch16	1040	3	51	(12,0) (14.6)	CHIANTI ([152])
ch16	1041	1	50	(14,0)	CHIANTI ([152,100])
ch17	1042	т 1	50	(10,0)	CHIANTI ([150])
ch17	1045	2	51	(10,0)	CHIANTI ([155] 157])
ch18	1044	3	51	(10,0)	CHIANTI ([155,157])
ch18	1045	4	50	(10,0) (20:6)	CHIANTI ([150])
ch10	1040	2	51	(20, 6)	CHIANTI ([150])
ch19	1047	4	50	(20,0) (12 14 16 18 20.7)	CHIANTI ([159,100])
ch 20	1040	+ /	50	(12, 14, 10, 10, 20, 7)	CHIANTI ([101])
ch20	1049	42	50	(20,7) (26.4)	CHIANTI ([102])
ch20	1050	3 1	51	(20,0) (14 16 19.9)	CHIANTI ([103,104])
ch21	1051	42	50	(14, 10, 10, 0) (9.7)	CHIANTI ([100])
CHZ1	1052	3	51	(0;7)	CHIAN II ([100])

 Table A2. Cont.

ID	Code	rt	dt	(Z; N)	References
ch22	1053	3	51	(12, 14, 16, 18, 20; 7)	CHIANTI ([161])
ch22	1054	4	50	(26;8)	CHIANTI ([167])
ch23	1055	3	51	(26;7)	CHIANTI ([161,162])
ch24	1056	4	50	(12, 14, 16, 20; 9)	CHIANTI ([168]), NIST
ch24	1057	3	51	(10, 12; 8)	CHIANTI ([169])
ch25	1058	3	51	(14, 16, 18; 8)	CHIANTI ([165])
ch25	1059	4	50	(26, 28; 9)	CHIANTI ([168,170,171]), NIST
ch26	1060	4	50	(14;10)	CHIANTI ([172])
ch26	1061	3	51	(26;8)	CHIANTI ([167])
ch27	1062	4	50	(16; 10)	CHIANTI ([173])
ch27	1063	3	51	(10;9)	CHIANTI ([174])
ch28	1064	3	51	(12;9)	CHIANTI ([174,175])
ch28	1065	4	50	(18,20,28;10)	CHIANTI ([173,176])
ch29	1066	4	50	(26:10)	CHIANTI ([177])
ch29	1067	3	51	(14:9)	CHIANTI ([174,178])
ch30	1068	3	51	(16;9)	CHIANTI $([174,179])$
ch30	1069	4	50	(12, 14, 16, 18, 20, 26, 28; 11)	CHIANTI $([125, 180])$
ch31	1070	4	50	(14:12)	CHIANTI, CHIANTI ([181])
ch31	1071	3	51	(20, 26; 9)	CHIANTI ([174,182])
ch32	1072	3	51	(14:10)	CHIANTI ([172])
ch32	1073	4	50	(16, 18, 20, 28; 12)	CHIANTI ([183]), NIST
ch33	1074	4	50	(14:13)	CHIANTL CHIANTI ([184])
ch33	1075	3	51	(16:10)	CHIANTI ([185])
ch34	1076	4	50	(16, 13)	(186 187)
ch34	1077	3	51	(18, 20, 28, 10)	CHIANTI ([176])
ch35	1078	3	51	(10,20,20,10) (26:10)	CHIANTI ([177])
ch35	1070	4	50	(26, 16) (26:13)	CHIANTI ([188])
ch36	1080	4	50	(26, 19)	CHIANTI ([189 190]) NIST
ch36	1081	3	51	$(12 \ 14 \ 16 \ 18 \ 20 \ 28.11)$	CHIANTI ([180])
ch37	1082	3	51	(14:12)	CHIANTI ([191])
ch37	1083	4	50	(26; 20)	CHIANTI ([192])
ch38	1084	3	51	$(16\ 18\ 20\ 28\cdot 12)$	CHIANTI ([183])
ch39	1085	3	51	(10, 10, 20, 20, 12) (14.13)	CHIANTI ([100])
ch40	1086	3	51	(11, 13)	CHIANTI ([186 187 194])
ch41	1000	3	51	(10, 10) (26, 18)	([100,107,171])
ch42	1007	3	51	(26, 10) (26: 19)	([190,190])
ch43	1000	3	51	(20, 1)	([100,100])
ci94	1002	3	56	(1:1)	[103 199]
cp99	1090	4	50	(26:21)	[200]
cp99	1091	3	56	(26, 21)	[200]
eg99	1092	3	56	(20, 21) (26:11)	[180 201]
ei99	1095	3	56	(20, 11) (26, 12)	CHIANTI [202]
fe01	1094	4	50	(20, 12) (26:25)	[203_206]
fo10	1095	3	56	(26, 23)	CHIANTI [207]
fo24	1090	3	56	(26, 17) (26, 3)	[127 208]
ar58	1097	4	50	(20,3)	[209]
im	1090	3	63	(7 12 16 18 28.1) (7.2)	Impact parameter method
1111 1/102	1100	4	50	(7, 12, 10, 10, 20, 1), (7, 2)	
km87	1100	3	56	(2,2)	[210]
kn80	1101	2	60	(10, 14, 10, 2) (6.2)	[10/]
kneah	1102	2	60	(0, 2) (2 8 12 20.2)	[104] [107 011]
kn802	1103	2	60	(2, 0, 12, 20, 2)	[104,107,211] [104,107,211]
KN09C	1104	5	69 E0	(20; 2)	[104-107,211]
mg05	1105	4	50	$(12; \delta)$ (10, 7)	[212], [213,214]
nouli	1106	4	50	(10;7)	CHIANTI, PP95 ([212,215])
11001	1107	A	EO	(10, 0)	OIII A MET DDOE (1010 0141)
ne02	1107	4	50 50	(10;8)	CHIANTI, PP95 ([212,214])

Table	A2.	Cont.

ID	Code	rt	dt	(Z;N)	Reference
ox01	1110	4	50	(8;6)	PP95 ([212,220]), CHIANTI ([149])
ox02	1111	4	50	(8;7)	CHIANTI, PP95 ([212,215])
ox03	1112	4	50	(8;8)	[218,219], PP95 ([212,214]
pb95	1113	3	56	(18, 20; 17)	[207]
pp01	1114	4	50	(6;6)	PP95 ([136,212,213])
pp01	1115	3	56	(6;6)	PP95 ([221–223])
pp02	1116	3	56	(7;7)	PP95 ([224,225])
pp03	1117	3	56	(8;8)	PP95 ([225,226])
pp04	1118	3	56	(16;14)	PP95 ([214])
pp05	1119	4	50	(18, 20; 16)	PP95 ([212,214])
pp05	1120	3	56	(18; 14), (20; 19)	PP95 ([214])
pp06	1121	3	56	(18;15)	PP95 ([227])
pp06	1122	4	50	(10;9), (20;17)	PP95 ([212])
pp07	1123	3	56	(18, 20; 16)	PP95 ([228])
pp07	1124	4	50	(16; 14)	PP95 ([212,229,230])
pp08	1125	4	50	(18;15)	PP95 ([212,231])
pp09	1126	4	50	(18; 17)	PP95 ([214])
pp10	1127	4	50	(20; 19)	PP95 ([232])
pp11	1128	4	50	(18; 14)	PP95 ([187,212])
rb96	1129	3	56 51	(16; 15)	[233], CHIANTI ([234])
sm00	1130	3	51	(20;13)	
5501	1121	4	50	(10, 13)	
5590 th02	1132	3	50	(10, 20, 15) (6 7 8 10 12 14 16 18 20 26.2)	[230]
7b96	1133	4	56	(0,7,6,10,12,14,10,10,20,20,2) (26,24)	[00]
Z1190 7097	1134	3	56	(20, 24) (26, 23)	[237]
zp77 7687	1136	3	56	(18, 28, 2)	[200]
2307 hm01	1130	3	56	(10, 20, 2) (26.3)	[100]
bm01	1138	4	50	(26:3)	[5]
bm02	1130	3	56	(26,3)	[8]
pm01	1140	4	50	(26, 7 - 10)	[6]
pm02	1141	4	50	$(26, 2^{-5})$ (26, 18-25)	[7]
mk01	1142	4	50	(26;10-17)	[9]
gm01	1143	4	50	(8:2-8)	[11]
gk01	1144	4	50	(7;1-7)	[17]
pm03	1145	4	50	(16; 2-16)	[14]
pq01	1146	4	50	(10, 12, 14, 16, 18, 20; 2-Z)	[15]
pq02	1147	4	50	(28;2-27)	[16]
pq03	1148	4	50	(14;2–13)	[22]
pq04	1149	4	50	(9, 11, 15, 17, 19; 2 - (Z - 1))	[26]
				(21-25, 27, 29, 30; 2-(Z-1))	
mb01	1150	3	56	(9, 11, 15, 17, 19, 21 - 25, 27; 3 - (Z - 2)),	Unpublished
				(28;19–26), (29,30;3–26)	
fj01	1151	3	56	(26; 2, 3)	[110]
fj01	1152	4	50	(26;2,3)	[110]
gu01	1153	4	50	(26; 11-21)	[109]
gu01	1154	13	6	(26; 11-21)	[109]
kf01	1155	5	25	(3-5,21,29,30;2),	[98]
				(6-8, 16; (Z-3)-Z), (10, 12, 14, 16, 18, (Z, 2), Z)	
				(10, 12, 14, 10, 10; (Z-2)-Z), (26, 28; (Z-2)-Z),	
				(20, 20, (Z - Z) - Z), (20, (Z - 1) - Z)	
bb01	1156	5 15	95	(20, (2 - 1) - 2) $(1 - 30 \cdot 1 - 7)$	[111]
th01	1157	5,15 7	53	(1-30, 1-2) (1-30, 1) $(21-25, 27-30, 2)$	[¹¹¹] Hydrogenic
1001	1107	1	00	(28:3-10,21-26,28)	i i y ai ogei ne
tb02	1158	7	53	(2-20.26:2)	[239]
tb03	1159	7	53	(6,7,8,10,12,14,16,18,20,26;3)	[240]
tb04	1160	7	53	(6,7,8,10,12,14,16,18,20,26;4)	[241]
				(, , , , , , , , , , , , , , , , , , ,	L

Table A	2. Cont.
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ID	Code	rt	dt	(Z;N)	Reference
tb05	1161	7	53	(6, 7, 8, 10, 12, 14, 16, 18, 20, 26; 5)	[242]
tb06	1162	7	53	(6,7,8,10,12,14,16,18,20,26;6)	[243]
tb07	1163	7	53	(7, 8, 10, 12, 14, 16, 18, 20, 26; 7)	[60]
tb08	1164	7	53	(8, 10, 12, 14, 16, 18, 20, 26; 8)	[60]
tb09	1165	7	53	(10, 12, 14, 16, 18, 20, 26; 9)	[60]
tb10	1166	7	53	(10, 12, 14, 16, 18, 20, 26; 10)	[244]
tb11	1167	7	53	(14, 18, 20, 26; 11)	[60]
tb12	1168	7	53	(26;12)	[245]
tb13	1169	7	53	(26;13)	[246]
tb14	1170	7	53	(14, 26; 14)	[60,247]
tb15	1171	7	53	(16;16), (18;18), (26;15–21)	[60]
tb16	1172	7	53	(26;22)	[248]
tb17	1173	7	53	(26;23)	[249]
tb18	1174	7	53	(26;24)	[250]
tb19	1175	7	53	(26;25)	[251]
tb20	1176	7	53	(26;26)	[252]
tb21	1177	7	53	(28;27)	[218]
vy01	1178	1,7	59	(3, 4, 5, 9, 11, 15, 17, 19, 21 - 25; 1 - Z),	[102]
				(27, 29, 30; 1-Z), (13; 1, 2, 12, 13),	
				(14;14), (16;16), (18;18),	
.		_		(20; 18, 19, 20), (26; 19–25), (28; 21–28)	
kp01	1179	7	53	(26;3–26)	[3,5,8,10]
gg01	1180	7	53	(6;3–6)	[21,37]
gm01	1181	7	53	(8;3-8)	[11]
gg02	1182	7	49	(8;6-8)	[23,32,33]
gk01	1183	7	53	(7;3–7)	[17]
wk01	1184	7	49	(10, 12, 14, 16, 18, 20; 3-10)	[18]
gg03	1185	7	49	(10; 8-10)	[13,35,253]
wk02	1186	7	49	(12, 14, 16, 18, 20; 11-17)	[19]
gg04	1187	7	49	(12; 3-12)	[25]
wk03	1188	7	49	(13; 3-11)	[24]
wk04	1189	7	49	(28;3-20)	[20]
tr01	1190	7	49	(9,11;3-(Z-2)), (15,17,19;3-11)	[27]
	1101	-	10	(21-25, 27, 29, 30; 3-11)	
tr02	1191	7	49	(15, 17, 19; 12 - (Z - 2)),	[28]
1 02	1100	-	10	(21-25, 27, 29, 30; 12-18)	
tr03	1192	7	49	(21-25, 27; 19-(Z-2)),	[29]
01	1100	0	1	(28; 21-26), (29, 30; 19-26)	
ap01	1193	8	1	(3-30;2), (15-30;14-Z)	[89,97]
ap01	1194	8	1	(3-5;2-2), (7,21,29,30;2),	[89,97]
- 101	1105	0	20	(13-25, 27-30; 13-2)	
ad01	1195	ð	38 20	(3-12; 3-2), (13-30; 3-13)	AMDDP([60])
adui	1196	ð	39	(3-13; 3-2), (14-30; 3-12), (26, 29, 12, 20), (26, 21, 26)	AMDPP ([64])
mc01	1107	0	22	(20, 28; 13-20), (26; 21-26) (12, 12), (14, 12, 14)	[00]
ns01	119/	ð	22	(13; 13), (14; 13, 14) (1, 20; 1)	[99] [100]
gtUI	1198	ð	30	(1-30;1)	[100]

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