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# Astronomy Astrophysics

# Electron impact excitation of Ar XVII\*

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**Abstract.** Energies for the lowest 49 levels among the  $1s^2$  and  $1sn\ell$  (n = 2-5) configurations of Ar XVII have been calculated using the GRASP code of Dyall et al. (1989, Comput. Phys. Comm., 55, 424). Additionally, radiative rates, oscillator strengths, and line strengths are calculated for all electric dipole (E1), magnetic dipole (M1), electric quadrupole (E2), and magnetic quadrupole (M2) transitions among these levels. Furthermore, collision strengths have also been calculated for all the 1176 transitions among the above 49 levels using the Dirac Atomic *R*-matrix Code (DARC) of Norrington & Grant (2005, Comput. Phys. Commun., in preparation), over a wide energy range up to 580 Ryd. Resonances have been resolved in the threshold region, and effective collision strengths have been obtained over a wide temperature range up to log  $T_e = 7.2$  K. Comparisons are made with the limited results available in the literature, and the accuracy of the data is assessed. Our energy levels are estimated to be accurate to better than 0.1%, whereas results for other parameters are probably accurate to better than 20%.

Key words. atomic data - atomic processes

# 1. Introduction

In two recent papers, we (Aggarwal et al. 2005; McKeown et al. 2004) have reported calculations of energy levels and radiative rates for transitions in Ar XIII – Ar XVI, and in this work we report similar results for transitions in Ar XVII. Additionally, we also present our calculations for collision strengths ( $\Omega$ ) and effective collision strengths ( $\Upsilon$ ) for transitions among the lowest 49 levels of the 1s<sup>2</sup>, 1s2s, 1s2p, 1s3 $\ell$ , 1s4 $\ell$  and 1s5 $\ell$  configurations.

Emission lines from the spectra of highly ionized argon have been observed in the UV, EUV and X-ray spectra of solar, stellar and other astrophysical plasmas by many space missions, such as SOHO, *Chandra* and XMM *Newton*. Some of the observed lines from the spectra of Ar XIV – XVIII have been specifically listed by Dere et al. (2001), and a complete list of lines over a wide range of wavelengths for many ions of argon are available in the CHIANTI database at http://wwwsolar.nrl.navy.mil/chianti.html.Emission lines of He-like ions, including those from Ar XVII, are also a prominent feature of the X-ray spectra of tokamak plasmas (Keenan et al. 1987; Phillips et al. 1994), which arise from impurity elements.

An analysis of observed spectra provides information on the temperature, density and chemical composition of the plasma. However, such an analysis requires information for a wide range of atomic parameters, including energy levels, radiative rates, and excitation rate coefficients. Since there is a paucity of measurements for these atomic parameters, theoretical results are of vital importance. A few calculations have been performed in the past by some authors, but most of these are limited to a few energy levels and/or transitions. Therefore, in this work we attempt to report results for a wider range of energy levels, and hence for a larger number of transitions. Furthermore, most of the available theoretical data are confined to the radiative rates for allowed and inter-combination (E1) transitions alone, whereas we here report similar data for other types of transitions as well, namely electric quadrupole (E2), magnetic dipole (M1) and magnetic quadrupole (M2), because these data are also required in the analysis and modelling of plasmas.

To report results for above named atomic parameters, we have employed the fully relativistic GRASP (General purpose Relativistic Atomic Structure Package) code of Dyall et al. (1989) for the generation of wavefunctions, and the Dirac Atomic *R*-matrix Code (DARC) of Norrington & Grant (2005) for the computations of  $\Omega$ , and subsequently of  $\Upsilon$ .

### 2. Energy levels

The  $1s^2$ , 1s2s, 1s2p,  $1s3\ell$ ,  $1s4\ell$  and  $1s5\ell$  configurations of Ar XVII give rise to 49 fine-structure levels, listed in Table 1. Our calculated energies obtained from the GRASP code, with and without including the Breit and QED effects, are given in this table along with those from the CHI-ANTI database and the experimental compilations of NIST

<sup>\*</sup> Tables 2-6 are available only in electronic form at the CDS via anonymous ftp to cdsarc.u-strasbg.fr (130.79.128.5) or via http://cdsweb.u-strasbg.fr/cgi-bin/qcat?J/A+A/441/831

(http://physics.nist.gov/PhysRefData). For our calculations, we have used the option of *extended average level* (EAL), in which a weighted (proportional to 2j + 1) trace of the Hamiltonian matrix is minimized. This produces a compromise set of orbitals describing closely lying states with moderate accuracy. Our calculations are in the jj coupling and configuration interaction (CI) has been included among the above stated 15 configurations. The inclusion of Breit and QED effects not only *lowers* the energies by a maximum of 0.3 Ryd (see, for example, levels 33–49), but also (slightly) alters the orderings in a few instances, such as for levels 21–24. However, we have retained the original orderings of the Coulomb energies, because these are the ones adopted in the subsequent tables.

It is clear from Table 1 that experimental energies are not available for all the levels, but the agreement with our results is within 0.2 Ryd (better than 0.1%), which is highly satisfactory. The energy levels of the CHIANTI database are from the calculations of Sampson et al. (1983). These results are slightly higher than our values (last column) or those of NIST, but agree closely with our data obtained without the inclusion of the Breit and QED effects. Additionally, energies of Sampson et al. are non-degenerate for many levels, such as for levels 37-49. However, Sampson et al. have clearly stated that their calculations are primarily for generating large quantities of collision strengths, and are not for accurate determination of transition energies, i.e. energy levels. For the same reason, they have estimated the accuracy of their energy levels to be within 1%, which is seen to be the case from Table 1. To conclude, we can state with confidence that our calculations have improved upon the energy levels of Sampson et al., which have also been beneficial in determining the level orderings.

#### 3. Radiative rates

The absorption oscillator strength  $(f_{ij})$  and radiative rate  $A_{ji}$  (in s<sup>-1</sup>) for a transition  $i \rightarrow j$  are related by the following expression:

$$f_{ij} = \frac{mc}{8\pi^2 e^2} \lambda_{ji}^2 \frac{\omega_j}{\omega_i} A_{ji} = 1.49 \times 10^{-16} \lambda_{ji}^2 (\omega_j / \omega_i) A_{ji}$$
(1)

where *m* and *e* are the electron mass and charge, respectively, *c* is the velocity of light,  $\lambda_{ji}$  is the transition energy/wavelength in Å, and  $\omega_i$  and  $\omega_j$  are the statistical weights of the lower *i* and upper *j* levels, respectively. Similarly, the oscillator strength  $f_{ij}$  (dimensionless) and the line strength *S* (in atomic unit, 1 au =  $6.460 \times 10^{-36}$  cm<sup>2</sup> esu<sup>2</sup>) are related by the following standard equations:

For the electric dipole (E1) transitions:

$$A_{ji} = \frac{2.0261 \times 10^{18}}{\omega_j \lambda_{ii}^3} S^{E1} \quad \text{and} \quad f_{ij} = \frac{303.75}{\lambda_{ji} \omega_i} S^{E1},$$
(2)

for the magnetic dipole (M1) transitions:

$$A_{ji} = \frac{2.6974 \times 10^{13}}{\omega_j \lambda_{ji}^3} S^{M1} \quad \text{and} \quad f_{ij} = \frac{4.044 \times 10^{-3}}{\lambda_{ji} \omega_i} S^{M1}, (3)$$

for the electric quadrupole (E2) transitions:

$$A_{ji} = \frac{1.1199 \times 10^{18}}{\omega_j \lambda_{ji}^5} S^{E2} \quad \text{and} \quad f_{ij} = \frac{167.89}{\lambda_{ji}^3 \omega_i} S^{E2}, \tag{4}$$

and for the magnetic quadrupole (M2) transitions:

$$A_{ji} = \frac{1.4910 \times 10^{13}}{\omega_j \lambda_{ji}^5} S^{M2} \quad \text{and} \quad f_{ij} = \frac{2.236 \times 10^{-3}}{\lambda_{ji}^3 \omega_i} S^{M2}.$$
(5)

In Table 2 we present transition energies ( $\Delta E_{ij}$  in Å), radiative rates ( $A_{ji}$  in s<sup>-1</sup>), oscillator strengths ( $f_{ij}$ , dimensionless), and line strengths (S in au), in length form only, for all 336 electric dipole (E1) and 391 electric quadrupole (E2) transitions among the 49 levels of Ar XVII. The indices used to represent the lower and upper levels of a transition have already been defined in Table 1. Similar results for 316 magnetic dipole (M1) and 410 magnetic quadrupole (M2) transitions are listed in Table 3. These results not only cover a wider range of transitions among larger number of levels than hitherto available in the literature, but are also for *all* possible transitions.

In Table 4 we compare our *f*-values for a limited set of transitions with those from the CHIANTI database, which are mainly from the calculations of Sampson et al. (1983). In this table we also list the ratio of our *f*-values in the length and velocity forms, in order to assess the accuracy of our results. For most of the transitions, the two forms of *f*-values agree within 20%, which is quite satisfactory. However, for some transitions, such as 3-32, 4-32, 6-32 and 7-35, the two forms of *f*-values differ up to an order of magnitude, but their magnitudes are invariably small, i.e.  $f \sim 10^{-3}$ . Similarly, agreement with the CHIANTI *f*-values is within 20% for a majority of transitions, especially those with larger *f*-values. However, for some transitions, such as 3-27, 3-37 and 9-27, the two sets of f-values differ by an order of magnitude. These are comparatively stronger transitions, and  $f_L/f_V$  is close to unity, which gives us confidence in our results. In general our results are higher with a few exceptions, such as for 6-13, 6-37 and 12-27 transitions, for which our *f*-values are lower by over an order of magnitude. However these transitions are comparatively weaker and  $f_L/f_V$  is close to unity. Based on this comparison of our results with the corresponding data from CHIANTI, and between  $f_L$  and  $f_V$ , we may state that for strong transitions  $(f \ge 0.01)$  our radiative rates are accurate to ~20%, but scope remains for improvement for the weaker transitions.

#### 4. Collision strengths

For the computations of collision strengths, we have employed the DARC program of Norrington & Grant (2005). This program includes the relativistic effects in a systematic way, in both the target description and the scattering model. It is based on the jj coupling scheme, and uses the Dirac-Coulomb Hamiltonian in the *R*-matrix approach. However, because of the inclusion of fine-structure in the definition of channel coupling, the matrix size of the Hamiltonian increases substantially. The *R*-matrix radius has been adopted to be 6.0 au, and 61 continuum orbitals have been included for each channel angular

Table 1. Target levels of Ar XVII and their threshold energies (in Ryd).

Index	Configuration	Level	Expt. <sup>a</sup>	CHIANTI <sup>b</sup>	<b>GRASP</b> <sup>c</sup>	<b>GRASP</b> <sup>d</sup>
1	$1s^2$	$\frac{1}{1}$ So	0.00000	0.000000	0.00000	0.00000
2	1\$2\$	${}^{3}S_{1}$	228.15015	228.225601	228.20039	227.97345
3	1s2p	$^{3}P_{0}^{\circ}$	229.52797	229.586395	229.55811	229.37311
4	1s2p	${}^{3}P_{1}^{\circ}$	229.57456	229.683594	229.64185	229.41945
5	1s2s	${}^{1}S_{0}$	229.64803	229.813187	229.79482	229.59423
6	1s2p	$^{3}P_{2}^{\circ}$	229.77757	229.942795	229.85488	229.61924
7	1s2p	${}^{1}\mathbf{P}_{1}^{\circ}$	230.75280	231.076797	230.91628	230.67177
8	1s3s	${}^{3}S_{1}$	270.03877	270.118805	270.10078	269.86520
9	1s3p	${}^{3}P_{0}^{\circ}$	270.41842	270.507599	270.48924	270.26436
10	1s3p	${}^{3}P_{1}^{\circ}$	270.43266	270.539978	270.51412	270.27814
11	1s3s	${}^{1}S_{0}$	270.43505	270.539978	270.56681	270.33115
12	1s3p	${}^{3}P_{2}^{\circ}$	270.49291	270.604797	270.57739	270.33735
13	1s3d	${}^{3}D_{1}$	270.70342	270.831604	270.76091	270.52312
14	1s3d	$^{3}D_{2}$	270.70332	270.831604	270.76537	270.52336
15	1s3d	$^{3}D_{3}$	270.72730	270.864014	270.78817	270.54629
16	1s3d	$^{1}D_{2}$	270.73840	270.896393	270.79950	270.55973
17	1s3p	${}^{1}\mathbf{P_{1}^{\circ}}$	270.75700	270.896393	270.90193	270.65251
18	1s4s	${}^{3}S_{1}$	284.49700	284.569214	284.51374	284.33206
19	1s4p	${}^{3}P_{0}^{\circ}$	284.65550	284.731201	284.66948	284.51012
20	1s4p	${}^{3}P_{1}^{\circ}$	284.66200	284.731201	284.67979	284.51602
21	1s4s	${}^{1}S_{0}$	284.66300	284.731201	284.70448	284.54750
22	1s4p	${}^{3}P_{2}^{\circ}$	284.68700	284.763611	284.76417	284.54187
23	1s4f	${}^{3}F_{2}^{\circ}$		284.893188	284.78987	284.59841
24	1s4f	${}^{3}F_{3}^{\circ}$		284.893188	284.79216	284.59761
25	1s4f	${}^{3}F_{4}^{\circ}$		284.893188	284.79899	284.60335
26	1s4f	${}^{1}F_{3}^{\circ}$		284.893188	284.80109	284.60404
27	1s4d	$^{3}D_{1}$	284.77650	284.860771	284.80115	284.61206
28	1s4d	$^{3}D_{2}$	284.77600	284.860771	284.80523	284.61259
29	1s4d	$^{3}D_{3}$	284.78659	284.893188	284.80532	284.62291
30	1s4d	$^{1}D_{2}$	284.79100	284.893188	284.80839	284.63107
31	1s4p	${}^{1}\mathbf{P}_{1}^{\circ}$	284.79900	284.893188	284.89237	284.69334
32	1s5s	${}^{3}S_{1}$	291.14600	291.211212	291.25271	290.94261
33	1s5p	${}^{3}P_{0}^{\circ}$	291.22781	291.276001	291.30708	291.02186
34	1s5p	$^{3}P_{1}^{\circ}$	291.23100	291.308411	291.32863	291.02504
35	1s5s	${}^{1}S_{0}$	291.23200	291.308411	291.32863	291.02873
36	1s5p	${}^{3}P_{2}^{\circ}$	291.24393	291.308411	291.33041	291.03802
37	1s5d	${}^{3}D_{1}$		291.373199	291.33041	291.08109
38	1s5d	$^{3}D_{2}$		291.373199	291.35444	291.08138
39	1s5d	$^{3}D_{3}$		291.373199	291.35978	291.08625
40	1s5f	${}^{3}F_{2}^{\circ}$		291.373199	291.37420	291.08943
41	1s5f	${}^{3}F_{3}^{\circ}$		291.373199	291.37423	291.08909
42	1s5d	${}^{1}D_{2}$	291.29700	291.373199	291.37860	291.08971
43	1s5f	${}^{3}F_{4}{}^{\circ}$		291.373199	291.37947	291.09206
44	1s5f	${}^{1}F_{3}^{\circ}$		291.373199	291.37950	291.09237
45	1s5g	${}^{3}G_{3}$		291.373199	291.40201	291.09235
46	1s5g	${}^{3}G_{4}$		291.373199	291.40291	291.09214
47	1s5g	${}^{3}G_{5}$		291.373199	291.40919	291.09395
48	1s5g	${}^{1}G_{4}$		291.373199	291.41330	291.09409
49	1s5p	${}^{1}P_{1}^{\circ}$	291.30100	291.373199	291.41497	291.09523

<sup>a</sup> NIST (http://physics.nist/gov/PhysRefData).
<sup>b</sup> Energy levels of Sampson et al. (1983) from CHIANTI database (http://wwwsolar.nrl.navy.mil/chianti.html).
<sup>c</sup> Present results from GRASP *without* Breit and QED effects.

<sup>d</sup> Present results from GRASP with Breit and QED effects.



**Fig. 1.** Partial collision strengths for the  $1s2s {}^{3}S_{1}-1s2p {}^{3}P_{1}^{\circ}$  (2–4) transition of Ar XVII, at three energies of 300 Ryd (circles), 400 Ryd (triangles), and 500 Ryd (stars).



**Fig. 2.** Partial collision strengths for the  $1s2p \, {}^{3}P_{1}^{\circ}-1s2p \, {}^{3}P_{2}^{\circ}$  (4–6) transition of Ar XVII, at three energies of 300 Ryd (circles), 400 Ryd (triangles), and 500 Ryd (stars).

momentum for the expansion of the wavefunction. This allows us to compute  $\Omega$  up to an energy of 580 Ryd. The maximum number of channels for a partial wave is 217, and the corresponding size of the Hamiltonian matrix is 13 293. In order to obtain convergence of  $\Omega$  for all transitions and at all energies, we have included all partial waves with angular momentum  $J \leq 39.5$ , although a higher range would have been preferable for the convergence of allowed transitions, especially those with  $\Delta n = 0$ . However, to account for the inclusion of higher neglected partial waves, we have included a top-up, based on the Coulomb-Bethe approximation for allowed transitions and geometric series for forbidden transitions.

In Figs. 1–3 we show the variation of  $\Omega$  with angular momentum *J* at three energies of 300, 400 and 500 Ryd, and for three transitions, namely 2–4 (1s2s  ${}^{3}S_{1}$ –1s2p  ${}^{3}P_{1}^{\circ}$ ), 4–6 (1s2p  ${}^{3}P_{1}^{\circ}$ –1s2p  ${}^{3}P_{2}^{\circ}$ ) and 4–14 (1s2p  ${}^{3}P_{1}^{\circ}$ –1s3d  ${}^{3}D_{2}$ ), which are "elastic" (i.e. allowed with  $\Delta n = 0$ ), parity



**Fig. 3.** Partial collision strengths for the  $1s2p \ ^{3}P_{1}^{\circ}-1s3d \ ^{3}D_{2}$  (4–14) transition of Ar XVII, at three energies of 300 Ryd (circles), 400 Ryd (triangles), and 500 Ryd (stars).

forbidden, and allowed ( $\Delta n \neq 0$ ), respectively. For the forbidden and allowed transitions shown in Figs. 2 and 3,  $\Omega$  have fully converged at all energies, including the highest energy of our calculations. However, for the "elastic" transitions our range of partial waves in not sufficient for the convergence of  $\Omega$ , as shown in Fig. 1. For such transitions a top-up from the Coulomb-Bethe approximation is quite significant.

In Table 5 we present our results of  $\Omega$  for all transitions in a wider energy range ( $300 \le E \le 550$  Ryd), but above thresholds. The indices adopted to represent a transition are already given in Table 1. These results of  $\Omega$  are not directly applicable in any diagnostic or modelling work, but are very useful in assessing the accuracy of a calculation, and will be helpful for future comparisons.

# 5. Effective collision strengths

Effective collision strengths  $\Upsilon$  are obtained after integrating  $\Omega$  over a Maxwellian distribution of electron velocities, i.e.

$$\Upsilon(T_{\rm e}) = \int_0^\infty \Omega(E) \exp(-E_j/kT_{\rm e}) d(E_j/kT_{\rm e})$$
(6)

where  $E_j$  is the incident energy of the electron with respect to the final state of the transition, k is Boltzmann's constant, and  $T_e$  is the electron temperature in K. Once the value of  $\Upsilon$ is known for a transition, the corresponding value of the excitation q(i, j) and de-excitation q(j, i) rate coefficients can be easily obtained from the following simple relations:

$$q(i,j) = \frac{8.63 \times 10^{-6}}{\omega_i T_{\rm e}^{1/2}} \Upsilon \exp(-E_{ij}/kT_{\rm e}) \qquad \text{cm}^3 \,\text{s}^{-1} \tag{7}$$

and

$$q(j,i) = \frac{8.63 \times 10^{-6}}{\omega_j T_{\rm e}^{1/2}} \Upsilon \qquad \text{cm}^3 \,\text{s}^{-1},\tag{8}$$

where  $\omega_i$  and  $\omega_j$  are the statistical weights of the initial (*i*) and final (*j*) states, respectively, and  $E_{ij}$  is the transition energy.

Since the threshold energy region is dominated by numerous resonances,  $\Omega$  have been computed at a large number of energies in order to delineate these resonances. We have performed our calculations of  $\Omega$  at ~13 000 energies in the threshold region. Close to thresholds (~0.1 Ryd above a threshold) the energy mesh is 0.001 Ryd, and away from thresholds is 0.002 Ryd. Thus care has been taken to include as many resonances as possible, and with as fine a resolution as is computationally feasible. However, the energy gap between the n = 2and 3 levels is very wide, i.e. ~40 Ryd – see Table 1. Therefore, in this energy region the mesh has been gradually increased to 0.01 Ryd.

The values of  $\Upsilon$  so computed are listed in Table 6 at a series of electron temperatures in the range 5.2  $\leq \log T_e \leq 7.2$  K, which is fully sufficient for the application of the data to solar, astrophysical and fusion plasmas. In earlier work, Zhang & Sampson (1987) have reported results of  $\Upsilon$  for transitions among the lowest seven levels listed in Table 1. In their calculations, they have adopted the Coulomb-Born-Exchange method and have also included the contribution of resonances in an approximate way - see Zhang & Sampson for details and discussion. Therefore, in Table 7 we compare our results of  $\Upsilon$ with theirs at two common temperatures of  $2.33 \times 10^6$  and  $1.17 \times 10^7$  K. Generally, the agreement between the two sets of results is within 20%, which is highly satisfactory. However, for the 1–2 (1s<sup>2</sup>  ${}^{1}S_{0}$ –1s2s  ${}^{3}S_{1}$ ) transition, our values of  $\Upsilon$  are higher by up to 50%, particularly towards the lower end of the temperature range. For this transition, we show our resonances in Fig. 4a in the entire threshold range. It is clear from this figure that there are dense and high resonances very close to the threshold, which have resulted in significantly higher values of  $\Upsilon$  at lower temperatures. In Fig. 4b we highlight these resonances in a narrow energy range below 231 Ryd. Since our calculations are in *jj* coupling (i.e. state levels are degenerate) in comparison to the intermediate coupling adopted by Zhang & Sampson, resonances are larger in our work. As a result of this, our values of  $\Upsilon$  are generally higher, although there are some exceptions, such as for the 4-5 transition.

The other results of  $\Upsilon$  available in the literature are by Keenan et al. (1987), who have interpolated results for transitions in Ar XVII from calculations for other He-like ions with nuclear charge  $8 \le Z \le 26$ . To be precise, they have adopted the R-matrix results of Kingston & Tayal (1984) and Tayal & Kingston (1984a,b, 1985) for CV, OVII and MgXI, and the distorted-wave results of Pradhan (1985) for CaXIX and Fe XXV. In Table 8 we compare our calculated results with their interpolated ones. For the lowest six transitions, the agreement between the two sets of Y values is remarkably good, and within 10%. However, for the remaining six transitions the discrepancy is up to a factor of two, and the present results are invariably higher. This is mainly because of our calculations being in the *jj* coupling scheme, which gives rise to more accurate determination of resonances, especially at energies close to thresholds. As an example, we demonstrate our resonances for the  $1s^2$   ${}^1S_0-1s3s$   ${}^3S_1$  (1–8) transition in Fig. 5, which clearly shows a few dense resonances close to the threshold. Considering that the earlier calculations, from which Keenan et al. interpolated their results, were in LS coupling, involved



**Fig. 4.** Collision strengths for the  $1s^{2-1}S_0-1s2s^{-3}S_1$  (1–2) transition of Ar XVII.



Fig. 5. Collision strengths for the  $1s^2$   $^1S_0$ -1s3s  $^3S_1$  (1–8) transition of Ar XVII.

**Table 7.** Comparison of effective collision strengths (Y) for transitions among the lowest 7 levels of Ar XVII.  $(a \pm b \equiv a \times 10^{\pm b})$ . First row: Zhang & Sampson (1987), Second row: present results.

Transition		Temperature (K)		Tra	ansition	Temperature (K)	
i	j	2.33+6	1.17+7	i	j	2.33+6	1.17+7
1	2	9.78-4	7.23-4	3	4	2.49-2	1.55-2
1	2	1.48-3	8.98-4	3	4	2.69-2	1.72-2
1	3	5.07-4	3.73-4	3	5	2.10-3	1.17-3
1	3	5.59-4	4.14-4	3	5	2.60-3	1.48-3
1	4	1.57-3	1.20-3	3	6	2.63-2	1.99-2
1	4	1.79-3	1.35-3	3	6	2.79-2	2.18-2
1	5	1.64-3	1.75-3	3	7	1.34-2	7.15-3
1	5	1.83-3	1.89-3	3	7	1.44-2	8.21-3
1	6	2.53-3	1.86-3	4	5	1.89-2	2.14-2
1	6	2.70-3	2.06-3	4	5	1.48-2	1.78-2
1	7	5.31-3	7.15-3	4	6	9.17-2	6.40-2
1	7	5.34-3	7.25-3	4	6	9.88-2	7.14-2
2	3	2.09-1	2.67-1	4	7	3.88-2	2.16-2
2	3	1.88-1	3.04-1	4	7	4.24-2	2.52-2
2	4	6.09-1	7.78-1	5	6	1.06-2	5.86-3
2	4	5.54-1	8.90-1	5	6	1.35-2	7.56-3
2	5	9.22-3	5.80-3	5	7	6.17-1	7.98-1
2	5	9.96-3	6.24-3	5	7	5.72-1	9.12-1
2	6	9.96-1	1.26-0	6	7	6.15-2	3.46-2
2	6	9.21-1	1.49-0	6	7	6.82-2	4.12-2
2	7	2.72-2	1.99-2				
2	7	2.83-2	2.36-2				

**Table 8.** Comparison of effective collision strengths ( $\Upsilon$ ) for some transitions of Ar XVII. ( $a \pm b \equiv a \times 10^{\pm b}$ ).

		Pı	esent Resu	lts	Keenan et al. (1987)				
Transition		Temperature (log, K)							
i	j	6.80	7.00	7.20	6.80	7.00	7.20		
1	2	1.118-3	9.531-4	7.946-4	9.454-4	8.319-4	7.184-4		
1	3	4.873-4	4.345-4	3.729-4	4.385-4	3.931-4	3.444-4		
1	4	1.568-3	1.408-3	1.224-3	1.356-3	1.246-3	1.131-3		
1	5	1.847-3	1.876-3	1.886-3	1.641-3	1.673-3	1.724-3		
1	6	2.417-3	2.157-3	1.847-3	2.191-3	1.963-3	1.719-3		
1	7	6.271-3	6.979-3	7.698-3	6.053-3	6.694-3	7.655-3		
1	8	2.717-4	2.352-4	1.995-4	1.428-4	1.385-4	1.307-4		
1	9	1.306-4	1.174-4	1.016-4	7.931-5	7.679-5	7.179-5		
1	10	4.002-4	3.625-4	3.173-4	2.527-4	2.484-4	2.379-4		
1	11	3.803-4	3.854-4	3.841-4	2.193-4	2.462-4	2.732-4		
1	12	6.360-4	5.715-4	4.945-4	2.917-4	3.081-4	3.068-4		
1	13	6.430-5	5.353-5	4.327-5					
1	14	1.086-4	9.390-5	8.029-5					
1	15	1.435-4	1.198-4	9.700-5					
1	16	1.173-4	1.119-4	1.086-4					
1	17	1.270-3	1.404-3	1.518-3	9.962-4	1.143-3	1.345-3		

limited CI and energy levels (and hence less resonances), and economised on the number of partial waves included, the agreement between the two sets of results in Table 8 is perhaps more than satisfactory, but is more qualitative than quantitative.

# 6. Conclusions

In the present work, results for energy levels, radiative rates, collision strengths, and effective collision strengths for transitions among the lowest 49 levels of Ar XVII have been presented for all transitions. Additionally, results for radiative rates have been presented for four types of transitions, namely E1, E2, M1 and M2. The present set of results cover a wider range of transitions than hitherto available in the literature, and hence are likely to be useful for modelling and diagnostics of a variety of plasmas. Additionally, our calculations have been performed in the *jj* coupling scheme, CI and relativistic effects have been included while generating wavefunctions, and a large range of partial waves has been adopted in order to achieve convergence in  $\Omega$  values for a majority of transitions. Furthermore, resonances have been resolved in a fine energy mesh in order to improve upon the accuracy of the determined values of  $\Upsilon$ . Similarly,  $\Omega$  have been computed in a wider energy range up to 580 Ryd in order to determine values of  $\Upsilon$  up to a temperature of ~1.6×10<sup>7</sup> K. Based on comparisons made with the earlier available data for several atomic parameters, our energy levels are assessed to be accurate to  $\sim 0.1\%$ , whereas the accuracy of other parameters is probably  $\sim 20\%$ . As stated above, an overall improvement has been made, both in range and accuracy, over the existing results for a limited number of transitions. However, scope remains for further improvement mainly by extending the work to higher levels of the  $n \ge 6$  configurations. Not only similar results for transitions with these higher levels will be useful for modelling work, but the accuracy of the present reported results will also be enhanced due to the inclusion of larger CI and hence more resonances arising from those. Until these improvements can be made in future work, we hope the present results will be highly useful for plasma applications.

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