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Atomic data for opacity calculations: XX. Photoionization cross sections and oscillator strengths for Fe II

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Abstract. Large scale *ab initio* calculations for the radiative data of Fe II have been carried out in the close coupling (cc) approbation employing the it-matrix method and a target state expansion consisting of 83 *LS* terms of Fe III. All bound states of Fe II with $n \le 10$ and $l \le 7$ are considered. The results include 1301 bound states in *LS* coupling, oscillator strengths for 35 941 transitions among the bound *LS* states, and detailed photoionization cross sections for all bound states. Autoionizing resonances, as well as the coupling to excited core states, enhance the photoionization cross sections substantially. The calculations of oscillator strengths have been extended beyond the requirement of the Opacity Project to include a large number of fine structure transitions in Fe II, using an algebraic transformation of the *LS* coupled line strength and the observed energies. The present f- values compare favourably with available experimental values and the calculations by Kurucz. However, the present results differ considerably from earlier 16-state R-matrix calculations and the new radiative data yield Rosseland mean opacities that are 50% higher. Some special features in the monochromatic opacity spectra of Fe II are also noted.

1. Introduction

Under the auspices of the Opacity Project (OP; Seaton 1987) ab initio calculations for accurate atomic radiative data for essentially all astrophysically abundant atoms and ions have been carried out by international collaborators, as reported in previous papers in this ADOC (Atomic Data for Opacity Calculations) series. The present work involves large scale computations for Fe II in the close coupling approximation, employing the R-matrix method as adapted for the OP. With many closely spaced energy levels, Fe II is a complex atomic system of 25 electrons where the electron correlation effects play a very important role. Recently Sawey and Berrington (1992) have reported R-matrix calculations for a few iron ions. Their Fe II calculations employed a close coupling (cc) expansion including only the states dominated by the 3d⁶ ground configurations of the residual ion or the 'target' ion Fe III. As we show in the present, much more extended work, it is necessary to also include a large number of additional terms in the eigenfunction expansion, dominated especially by the excited 3d⁵4s and the 3d⁵4p configurations, in order to obtain accurate radiative parameters. The aim of the present R-matrix close coupling calculations of Fe II is to take into account all the important states of Fe II in the wavefunction expansion and obtain more accurate radiative data for energy levels, oscillator strengths and photoionization cross sections. The importance of such calculations and a summary of theoretical details can be obtained in the first two papers of the ADOC series (Seaton 1987, Berrington et al 1987). In an earlier brief report

(Le Dourneuf *et al* 1993), we reported the first detailed calculation, with autoionizing resonances and channel coupling effects, for the photoionization of the states with the ground state symmetry 6 D. The present report is a complete account of the comprehensive calculations including 90 total Fe II symmetries, and photoionization of all bound states up to $n \le 10$ and $l \le 7$ and oscillator strengths of all possible bound-bound transitions among the 1301 computed states.

In addition to the extensive LS coupling calculations for the OP calculations, we also obtain fine structure oscillator strengths for Fe II through an algebraic transformation of the LS multiple line strengths and using observed spectroscopic energies. Most of the resulting fine structure f-values compare favourably with available experimental data, and rather better than those calculated by Kurucz (1981) using semi-empirical methods†. At present the data set by Kurucz is the only available source of Fe II f-values for most applications where a large number of transitions need to be considered. The present approach yields f-values for all transitions between the fine structure components of the dipole allowed LS multiplets, given the observed energies of the individual components. Thus a reasonably complete data set for the large number of Fe II f-values is obtained.

2. Target states

Following the convention of collision theory, we refer to the ion in the e + ion system as the 'target' ion (also, as the 'core' or the 'residual' ion following photoionization). The importance of accurate target representation in cc calculations is to be emphasized as the necessary first step. For the cc calculations for Fe II we obtain the target eigenfunctions for Fe III using the SUPERSTRUCTURE program by Eissner et al (1974) based on a scaled Thomas-Fermi-Dirac potential and configuration interaction (CI) wave-functions. Given the number of states needed in the calculations, the atomic structure calculations are rather complicated since proper account needs to be taken of the CI effects, while the total number of configurations must be kept small to minimize, as much as possible, the memory of the CPU requirements. The task was found to be particularly difficult and time consuming for Fe III due to the large number of target states considered. The principal configurations (i.e. whose terms are explicitly included in the CC expansion) are: 3d⁶, 3d⁵4s and 3d⁵4p. It might be noted here that it is the dipole core transitions between the even and the odd parity configurations that give rise to the well known photoexcitation-of-core (PEC) resonances in photoionization cross sections (Yu and Seaton 1987, Nahar and Pradhan 1991). The PEC resonances were not considered in the earlier work by Sawey and Berrington (1992) since the excited configurations were not included. As the 3d shell plays the dominant role in electron correlation, the correlation configurations are constructed mainly with respect to variations in the 3d orbital and excitations involving the 3d electrons. In addition, important improvements in the accuracy of the Fe III energies and oscillator strengths were achieved by introducing a correlation configuration with the 4d orbital (which increased the optimization time considerably). The final configuration list and the

 $\dagger f$ -values given in this paper correspond to the most recent gf values calculated by him which were obtained from him by private communication.

Thomas-Fermi scaling parameters λ_{nl} are given in table 1. The choice of the target states actually included in the cc calculations was somewhat independent of the target optimization since all 136 *LS* terms dominated by the three principal configurations, $3d^6$, $3d^54s$ and $3d^54p$, are well represented with the choice of the target in table 1. We include up to 83 terms in the present Fe II calculations. In particular we include all odd parity terms dominated by $3d^54p$ that are linked via dipole transitions to the ground state 5D to take account of strong PEC resonances in the photoionization cross sections.

Table 1 compares the energies of the 83 terms of Fe III with the observed values (Sugar and Corliss 1985, Moore 1952). It may be noted that three calculated singlet terms: 3d⁶ (¹D, ¹S), 3d⁵ ²S4s ¹S, have not been observed. Comparison of the calculated target energies with the observed energies shows agreement within 10% for most of the states, the largest discrepancy being about 18% for the 3d⁶ ¹I state.

Over 300 dipole oscillator strengths were obtained from SUPERSTRUCTURE for the *LS* target states dominated by the 3d⁶, 3d⁵4s and 3d⁵4p configurations. These oscillator strengths show agreement between the length and the velocity forms within 15% for most of the transitions in Fe III, further confirming the overall accuracy of the large set of the target eigenfunctions.

In table 1, the notations O, S, Q and D in the target state column specify that the corresponding Fe III state couples to octet, sextet, quartet or doublet symmetry of the e + ion system, $SL\pi$, of Fe II respectively. Thus there are 2 O, 21 S, 58 Q and 62 D such terms of Fe III that are in the target expansion for radiative calculations of octet, sextet, quartet and doublet states of Fe II. In table 1, the number next to each notation of O, S, Q and D is the energy degeneracy number for that state. This will be explained in the following section.

3. Computations and calculations for the radiative data

As in the case of all OP work, the present computations have been carried out in LS coupling, that is, relativistic effects are not taken into account. Since Fe II is a singly charged ion, it has been assumed that LS coupling would provide a good approximation for the radiative data. All bound states, denoted as S_tL_tnl , where S_tL_t is a target state, and $n \le 10$ and $l \le 7$ are considered for the radiative data.

Each excited target state S_tL_t of the ion is the series limit for the Rydberg series S_tL_tvl of the (N+1) electron system, where v is the effective quantum number of the (N+1)th electron. These are pure bound states if they lie below the first ionization threshold, but those that lie above the first ionization threshold are usually quasibound states and manifest themselves through autoionizing resonances in the photoionization cross sections (some states above the first ionization threshold may be pure bound states in LS coupling if they are forbidden to autoionize into the corresponding continua). These Rydberg resonances repeat the pattern for each increment of v. As v increases, the resonances get narrower and numerical resolution becomes difficult. To obviate the problem, we employ a constant mesh in v, for each interval v and v+1, to fully delineate the Rydberg resonances up to v=10 with a mesh interval of $\Delta v=0.01$. The region $10 < v \le \infty$ that we term as the QDT region, corresponds to a small energy region which is treated through quantum defect theory (QDT) using the Gailitis averaging method (e.g. Nahar and Pradhan 1991).

For closely spaced target states, as in the case of Fe III, the QDT region of different target

Table 1. Calculated (cal) term energies of Fe III (configurations $3d^4$. $3d^54s$ and $3d^54p$) and comparison with the observed (obs) energies. The energies, in Ry, are relative to the $3d^{65}D$ ground state. The three states, $3d^54s$ 'S, $3d^6$ 'D and $3d^6$ 'S, are unobserved. The notation O, S, Q and D in the target state column specifies the coupling to the octet, sextet, quartet and doublet symmetries of Fe II respectively, The number next to them represents the degeneracy index (see text). The spectroscopic and correlation configurations for Fe III and the values of scaling parameter λ_{nl} for each orbital in the Thomas-Fermi potential are also given.

				Energy	•	
	State		Obs.	Calc.	Target	states
1	3d ⁶	⁵ D	0.0	0.0	[\$1	Q1]
2	3d ⁶	³ P2	0.1826	0.1810	[Q2	D1]
3	3d ⁶	³H	0.1845	0.2163	[Q3	D2]
4	3d ⁶	³F2	0.1972	0.2105	[Q3	D2]
5	$3d^6$,C	0.2263	0.2537	[Q3	D2]
6	3d ⁵ (⁶ S)4s	⁷ S	0.2742	0.2728	[01	S2]
7	3d ⁶	1	0.2766	0.3246	[D3]
8	3d ⁶	³D	0.2805	0.3037	[Q4	D3]
9	3 d ⁶	'G2	0.2815	0.3021	[D3]
10	3d ⁶	'S2	0.3172	0.3190	ĺ	D3]
11	3d ⁶	D2	0.3263	0.3460	E	D3]
12	3d ⁵ (⁶ S)4s	sS.	0.3736	0.4167	[\$3	Q5J
13	3d ⁶	¹F	0.3909	0.4295	Ĺ	D4]
14	3d ⁶	³ P1	0.4556	0.4900	[Q6	D5]
15	3d ⁶	3F1	0.4580	0.4985	[Q6	D5]
16	3d ⁶	¹G1	0.5214	0.5738	Ĩ	D6]
17	3d ⁵ (⁴ G)4s	⁵G	0.5783	0.5995	[S4	Q7]
18	3d ⁵ (4P)4s	⁵P	0.6061	0.6466	[\$5	Q8]
19	3d⁵(⁴D)4s	5D	0.6358	0.6730	[\$5	Q8]
20	3d ⁵ (⁴ G)4s	^{3}G	0.6444	0.6957	[Q9	D7]
21	3d ⁵ (⁴P)4s	³ P	0.6724	0.7438	[Q9	D7]
22	3d ⁵ (⁴ D)4s	^{3}D	0.7019	0.7691	[Q10	D8]
23	3d ⁵ (² I)4s	3 I	0.7276	0.7526	[Q10	D8]
24	3d ⁵ (⁶ S)4p	⁷ ₽°	0.7516	0.7338	[O2	S6]
25	$3d^6$	¹D		0.7650	[D8]
26	3d ⁵ (2D3)4s	^{3}D	0.7510	0.8126	[Q1!	D9]
27	3d ⁵ (⁴ F3)4s	5F	0.7585	0.8038	[56	QII]
28	3d ⁵ (² I)4s	Ţ	0.7603	0.8005	Ĺ	D9]
29	3d ⁵ (² F2)4s	³F	0.7689	0.8211	[Q11	D9]
30	3d ⁵ (² D3)4s	'D	0.7914	0.8601	ĺ	D10]
31	3d ⁵ (² F2)4s	¹F	0.8010	0.8693	[D10]
32	3d ⁵ (² H)4s	3H	0.8090	0.8524	[Q12	D10]
33	3d ⁵ (⁶ S)4p	5P°	0.8133	0.8295	[S7	Q12]
34	3d ⁵ (2G2)4s	³G	0.8184	0.8679	[Q13	D10]
35	3d ⁵ (⁴ F)4s	³F	0.8244	0.8991	[Q13	D11]
36	3d ⁵ (² H)4s	^¹ H	0.8431	0.9000	Ī	D11]
37	3d ⁵ (² F1)4s	³F	0.8511	0.9106	[Q13	D11]
38	3d ⁵ (2G2)4s	'G	0.8521	0.9168	1	D11)
39	3d ⁵ (² F1)4s	^t F	0.8843	0.9582	Ī	D12]
40	3d ⁶	¹S		0.9596	į	D12]
41	3d ⁵ (² S)4s	3S	0.8991	0.9721	[Q14	D12]
42	3d ⁵ (2D2)4s	³D	0.9652	1.0406	[Q15	D13j
43	3d5(2D2)4s	'D	0.9985	1.0908	i	D13j
44	3d ⁵ (⁴ G)4p	¹G°	1.0358	1.0353	[S8	Q15]
45	3d ⁵ (2S)4s	¹S		1.0310	į	D13]
46	3d ⁵ (2G1)4s	³G	1.0419	1.1137	[Q16	D14j
47	3₫ ⁵ (⁴G)4p	5H°	1.0512	1.0507	[S9	Q16]

Table 1. (continued)

				Energy		
	State		Obs.	Calc.	— Target	states
48	3d ⁵ (⁴ G)4p	⁵F°	1.0619	1.0717	[S9	Q16]
49	3d⁵(⁴P)4p	⁵S°	1.0653	1.0846	. [S9	Q16]
50	3d⁵(⁴P)4p	5D°	1.0661	1.0825	[89	Q16]
51	3d ⁵ (2G1)4s	'G	1.0748	1.1619	[D14]
52	3d⁵(⁴G)4p	3F°	1,0778	1.0965	[Q16	D14]
53	3d ⁵ (⁴ G)4p	3Ho	1.0800	1.0963	[Q16	D14]
54	3d ⁵ (⁴ P)4p	5Po	1,0810	1.1083	[89	Q16]
55	3d ⁵ (⁴ P)4p	³P°	1.0921	1.1257	[Q17	D14]
56	3d ⁵ (⁴ D)4p	5F°	1.1041	1.1220	[S10	Q17]
57	3d5(4G)4p	³G°	1.1112	1.1475	[Q17	D14]
58	3d ⁵ (⁴ P)4p	$^3\mathbf{D}^{\mathbf{o}}$	1.1167	1.1567	[Q17	D14]
59	3d ⁵ (⁴ D)4p	5D°	1.1208	1.1449	[\$10	Q17j
60	3d ⁵ (⁴ D)4p	5 p°	1.1272	1.1509	{S10	Q17]
61	3d ⁵ (⁴ D)4p	$^{3}D^{\circ}$	1.1381	1.1721	[Q18	D15]
62	3d⁵(⁴D)4p	³F°	1.1442	1.1784	[Q18	D15]
63	3d ⁵ (⁴ P)4p	3S°	1.1518	1.2158	[Q18	D15]
64	3d ⁵ (⁴ D)4p	³ P°	1.1733	1.2231	[Q18	D15]
65	3d ⁵ (² I)4p	3K°	1,1873	1.1884	[Q18	D15]
66	3d ⁵ (² l)4p	3Io	1.1912	1,1932	[Q18	D15]
67	3d ⁵ (² I)4p	'H°	1.2002	1.2104	Ī	D15]
68	3d ⁵ (² I)4p	'K°	1.2028	1.2102	Ē	D15]
69	3d5(2D)4p	³F°	1,2064	1.2815	[Q18	D15]
70	3d ⁵ (² I)4p	³H°	1.2072	1.2223	[Q18	D15]
71	$3d^{5}(^{2}D)4p$	¹D°	1.1978	1,2375	i	D15]
72	$3d^{5}(^{2}D)4p$	³P°	1.2252	1.2698	[Q18	D15]
73	3d ⁵ (² F)4p	¹G°	1.2244	1.2554	į i	D15]
74	3d ⁵ (² F)4p	$^3G^{\circ}$	1.2320	1.2635	Q18	D15]
75	3d ⁵ (⁴ F)4p	⁵G°	1.2339	1.2602	[S11	Q18]
76	$3d^5(^2D)4p$	$^{3}D^{\circ}$	1,2313	1.2790	[Q18	D15]
77	3d ⁵ (a ² F)4p	$^{3}D^{\circ}$	1.2413	1.2859	[Q18	DIS]
78	$3d^{5}(^{2}I)4p$	1 1 0	1.2369	1.2618	i	D15]
79	3d ⁵ (² D)4p	¹F°	1.2411	1.2741	i	D15]
80	3d ⁵ (⁴ F)4p	5F°	1.2402	1.2716	เรา	Q18]
81	$3d^5(a^2F)4p$	3F°	1.2453	1.2347	[Q18	D15]
82	3d ⁵ (⁴ F)4p	5D°	1.2520	1.2847	[S11	Q18
83	3d ⁵ (² H)4p	³H°	1.2566	1.2781	[Q18	D15

Fe III configurations:

Spectroscopic: 1s²2s²2p⁶3s²3p⁶3d⁶, 1s²2s²2p⁶3s²3p⁶3d⁵4s, 1s²2s²2p⁶3s²3p⁶3d⁵4p

Correlation: 1s²2s²2p⁶3s²3p⁴3d⁸, 1s²2s²2p⁶3p⁶3d⁸, 1s²2s²2p⁶3s3p⁶3d⁷,

1s²2s²2p⁶3s3p⁵3d⁸, 1s²2s²2p⁶3s²3p⁶3d⁵4d

 λ_{nl} : 1.1 (1s) 1.1 (2s), 1.1 (2p), 1.095 (3s), 1.091 (3p), 1.0341 (3d) 1.04351 (4s)

1.044 66 (4p), 1.278 65 (4d)

as degenerate in the present radiative calculations. We expect little consequent loss of accuracy since most of these states lie fairly high in energy. In table 1, the number of such terms that are treated degenerate is given next to the notation O, S, Q and D, mentioned earlier.

Following the R-matrix calculations for the e + Fe III system, we first obtain the energy levels of Fe II for all $SL\pi$ symmetries considered. For complex atoms and ions isoelectronic with the third and the fourth row elements, it is a non-trivial task to identify all the computed levels and a careful analysis is required based partly on a study of quantum defects along overlapping Rydberg series, and partly on the contributions of the closed channel wavefunctions, in the region outside the R-matrix boundary, to the total bound state. Nearly all Fe II states have been unambiguously identified. It might be noted that for the calculation of opacities the level identification problem is not consequential. However, we strive to attain precise LS term designations in order to facilitate other applications of the present data, particularly in the important extension of the OP work to obtain f-values for fine structure components within LS mutiplets.

The work on Fe II was carried out on the 8 processor 64 MW Cray Y-MP at the Ohio Supercomputer Center, Columbus, Ohio. Table 2 shows the total CPU time required for the radiative calculations for octet, sextet, quartet and doublet symmetries, and the maximum memory needed for the R-matrix close coupling calculations with 83 state expansion for Fe II. It required up to 20 MW of memory for the largest $SL\pi s$ and a total of about 450 CPU hours. Work was divided according to different symmetries, and the table shows the number of target states coupled to each $SL\pi$ and used as the target set of eigenfunctions for radiative calculations for that particular symmetry. For the largest of the quartet and doublet symmetries, the R-matrix calculations could be carried out for only one $SL\pi$ at a time.

Table 2. Summary of the radiative calculations for Fe II: $N_{\rm CC}$ is the number of target states coupled to a particular spin symmetry of Fe II, CPU is the amount of time required for that symmetry. Maximum memory requirement and disk space for a typical quartet or doublet symmetry run are given below. $N_{SL\pi}$ the total number of bound symmetries, $SL\pi$ shows the range of these symmetries and N_{E_T} is the corresponding number of bound states up to $n \le 10$, $l \le 7$. N_{bnd} is the number of bound states below the first ionization threshold, N_f is the number of oscillator strengths. The largest case is $SL\pi=2G$ with 181 continuum channels and the hamiltonian matrix size of 2228. Memory and disk space range requirements: RAM: 20 MW, disk: 3.5-4 GB.

Symmetry	Ncc	ÇPU	N _{SLx}	SLπ (range)	N _{E_T}	N_{bnd}	N_f
Octets	2	1.2 min	16	⁸ S- ⁸ L, ⁸ S ⁿ - ⁸ K°	92	6	95
Sextets	21	4.2 h	25	65-6N, 65°-60°	234	205	4 267
Quartets	58	166.5 h	29	⁴ S- ⁴ T, ⁴ S°- ⁴ R°	357	308	9 965
Doublets	62	174.6 h	20	${}^{2}S_{-}{}^{2}M, {}^{2}S_{-}{}^{2}M^{\circ}$	618	224	21 614.
Total	83	446 h†	90		1301	743	35 941

[†] Including about 100 CPU h spent on trials and problems

4. Results and discussions

Three sets of data are calculated: (a) energy levels, (b) oscillator strengths and (c) photoionization cross sections; these are discussed below with selected examples.

4.1. Energy levels

We obtain 1301 LS bound state and identify 743 states that lie below the first ionization threshold of Fe II, i.e. below the 3d⁶(5 D) ground state of Fe III. In addition, a few bound states are obtained that lie above the ionization threshold but are forbidden to autoionize in LS coupling. The number of bound states that have been identified are more than twice the number that have been reportedly observed. Table 2 gives a summary of the number of bound state symmetries $SL\pi$, their total L and S value ranges, and the corresponding number of bound states computed for each $SL\pi$. All of the observed LS terms, 266 in total, have been calculated and identified. This could not have been

possible with the earlier 16-cc *R*-matrix calculations (Savvey and Berrington 1992) because several of the observed states couple to terms dominated by excited configurations 3d⁵4s and 3d⁵4p of Fe III which are not included in the earlier work. The present calculated energy for the 3d⁶4s(⁶D) ground state of Fe II differs by 0.6% from the observed value, compared to a 7% discrepancy in the previous calculation.

In table 3 we compare the calculated and observed energies. The latter set includes recent measurements from the Lund group (Johansson 1992); the *LS* energies have been computed as the statistically weighted average over the fine structure components. In a small number of cases the set of observed fine structure levels is incomplete; such states are marked with asterisks. Comparison shows that most of the calculated *LS* term energies are within 10% of the observed ones, yet many do show larger differences of up to 10-30%. Exclusion of relativistic effects is probably the prime contributor to the discrepancies. While the relativistic calculations are planned, as part of a new project on the iron-peak elements (the Iron Project), using Breit-Pauli *R*-matrix method, it is estimated that the *ab initio* fine structure calculations may require an order of magnitude more resources and effort even over the present one.

4.2. Oscillator strengths

Dipole oscillator strengths (*f*-values) for approximately 36 000 transitions among the 1301 calculated bound states of Fe II are obtained in *LS* coupling. Over 19 000 of these transitions are between bound states which lie below the first ionization threshold. For opacity calculations we also include transitions of bound states when the lower state lies below the first ionization threshold and the upper state lies above; since the latter do not appear as resonances in the photoionization cross sections in *LS* coupling but the corresponding oscillator strength does contribute to total photoabsorption. Table 2 lists the number of oscillator strengths obtained for each spin symmetry and all corresponding total angular momenta *L*. Each oscillator strength in *LS* coupling corresponds to a number of transitions, when we consider the fine structure, resulting in over 100 000 individual *f*-values. These calculations are discussed below.

As an enormous amount of data have been computed, one of the primary aims of this report is to attempt to establish the uncertainties involved relative to available experimental data and previous theoretical calculations. Table 4(a) presents selected comparisons with other results found in literature. The present oscillator strengths are obtained from the calculated line strength (s), and the observed energies, according to the relation, $S = (3g_r/E_f)f_{if}$. Of the two sets of columns for transition of states in the table, the first set of columns compares the present results with both the measured values compiled by NIST (Fuhr *et al* 1988) and the calculated ones by Kurucz (1981), and the second set of columns with those of Kurucz. NIST has compiled and evaluated all the available measured and some theoretical values for the oscillator strengths. The column listing the NIST f-values for dipole allowed transitions in LS coupling are averaged over the fine structure transitions for most cases. Kurucz obtained the f-values using semi-empirical atomic structure calculations including some relativistic effects (his f-values quoted in table 4(a) are statistically averaged over the fine structure).

For most cases the present values agree within 10% with those by Kurucz. Overall we find that the present *LS* multiplet oscillator strengths are in somewhat better agreement with the experimental values than those of Kurucz for most transitions (this *is* also true of the fine structure *f*-values discussed in the next section).

Table 3. Comparison of calculated (cal) energies (in Ry) of the octet, sextet, quartet and doublet states of Fe II with the observed (obs) ones. * denotes that the observed *LS* energy is obtained from incomplete set of fine structure levels The table contains the most recent measured values of the energy levels at Lund (Johansson 1992).

			E (Ry)			E ((Ry)
State		Obs.	Calc.	State		Obs.	Calc.
			Octo	ets: 2-cc			
3d ⁵ ⁶ S4s4p ³ P°	z^8P^o	0.709 72	0.7783	3d ⁵ 4s ⁷ S 5s	⁸ S	0.250 23	0.2457
3d ⁵ 4p ²	8P	0.215 67	0.2742	3d ⁵ 4s ⁷ S 4d	8D	0.192 13	0.2300
			Sexte	ts: 21-cc			
3d ^{6 5} D4s	a ⁶ D	1.185 91	1.1782	3d ^{5 4} D4s4p	6D0	0.333 45	0.3478
3d ⁵ 4s ²	a ⁶ S	0.977 21	0.9951	3d ⁵ D4s4p	⁶ P°	0.327 51	0.3443
3d ^{5 5} D4p	z^6D^0	0.836 95	0.8466	3d ^{s s} D6s	۴D	0.259 37	0.2555
3d ^{5 5} D4p	z^6F^0	0.805 42	0.8177	3d ^{5 5} D5d	۴F	0.240 42*	0.2454
3d ^{5 5} D4p	z^6P^0	0.797 26	0.8111	3d⁵ ⁵D5d	"P	0.239 87	0.2438
3d ^{3 6} S4s4p	y ⁶ P°	0.623 92	0.6594	3d ^{5 5} D5d	°D	0.236 33	0.2422
3d ^{5 6} S4s4p	x ⁶ P°	0.467 09	0.4785	3d ⁵ 5D5d	6G	0.234 92	0.2417
3d⁵ ⁵D5s	e ⁶ D	0.476 40	0.4687	3d ^{5 5} D5d	⁶ S	0.226 38	0.2388
3d⁵ ⁵D4d	e ⁶ F	0.423 50	0.4312	$3d^54p^2$	۴D	0.218 62	0.2713
3d⁵ ⁵D4d	۴D	0.426 04	0.4273	3d ⁵ 4s ⁷ S5s	⁶ S	0.218 35	0.2117
3d ^{5 5} D4d	6₽	0.421 30	0.4219	3d ^{5 4} F4s4p	6F°	0.218 56	0.2275
3d ^{5 5} D4d	e ⁶ G	0.420 21	0.4223	3d ⁵ 5D6p	٥D٥	0.217 65	0.2197
3d ^{5 5} D4d	⁶ S	0.410 61	0.4128	3d ^{5 5} D6p	⁶ F°	0.212 07	0.2146
3d⁵ ⁴G4s4p	y°F°	0.392 40	0.3756	3d ^{5 5} D6p	6	0.210 58	0.2126
3d ⁵ ⁴P4s4p	6Do	0.385 57	0.4120	3d ⁵ 4F4s4p	6D₀	0.204 51	0.2116
3d ^{5 5} D5p	6D°	0.378 96	0.3789	3d ⁵ 5D7s	6D	0.162 27	0.1638
3d5 4P4s4p	6Po	0.375 53	0.3966	3d5 5D6d	6F	0.153 54*	0.1568
3d ⁵ ⁵ D5p	6Fo	0.367 66	0.3675	3d5 5D6d	⁵G	0.151 48*	0.1550
3d ^{5 5} D5p	⁶ P°	0.356 31	0.3581	$3d^54p^2$	6P	0.141 66	0.1834
3d ^{5 4} D4s4p	6F°	0.349 40	0.3756	3d 54s 7S4d	۴D	0.129 55	0.1446
•			Quart	ets: 58-cc			
3d ⁷	a ⁴ F	1.167 68	1.081	3d ⁵ 4s ³ I4p	۴K°	0.254 88	0.2852
3d ^{6 5} D4s	a⁴D	1.113 88	1.1060	3d ^{6 5} D6s	۲D	0.253 38	0.2531
3d ⁷	a ⁴ P	1,065 66	0.9846	3d ⁶ ³ G5s	f⁴G	0.252 41	0.2403
3d ^{6 3} P4s	b ⁴ P	0.994 50	0.9313	3d ⁵ 4s ³ 14p	4 _J °	0.250 27	0.2810
3d ^{6 3} H4s	a ⁴ H	0.994 15	0.9631	3d ⁶ ³ P4d	⁴D	0.249 70*	0.2206
3d ^{6 3} F4s	ь⁴F	0.981 86	0.9327	3d ⁶ ³ H4d	¹K	0.244 07*	0.2200
3d ⁶ ³ G4s	a⁴G	0.954 95	0.9327	3d ^{6 3} H4d	⁴H	0.244 07	0.2120
3d ⁶ ³ D4s	b ⁴ D	0.903 38	0.8499	3d ⁶ ³ H4d	⁴G	0.243 87	0.2104
3d ^{6 5} D4p	z ⁴ F°	0.782 23	0.7968	3d ^{6 5} D5d	⁴F	0.243 20	0.2112
3d ^{6 5} D4p	z⁴D°	0.781 97	0.7890	3d ⁶ ³ P4d	⁴F	0.242 56	0.2377
3d ^{6 5} D4p	z ⁴ P°	0.759 42	0.7706	3d ⁶ ³ H4d	4I	0.241 46	0,2106
3d ^{6 3} P4s	c ^e P	0.735 96	0.6515	3d 114d 3d ^{6 3} P4d	⁴P	0.234 41	0.2206
3d ^{6 3} F4s	c ⁴ F	0.732 69	0.6462	3d 54s 314p	⁴ H°	0.234 41	0.2206
3d ⁵ 4s ²	b ⁴ G	0.695 23	0.6913	3d 4s 14p 3d 4s D4p	⁴F°	0.234 39	0.2367
3d ⁵ 4s ²	d⁴P			3d 4s D4p 3d ^{6 5} D5d	⁴D		
3d ^{6 3} P4p		0.666 03	0.6331	3d ⁶ ³ H4d	⁴F	0.232 53*	0.2404
3d*4s ²	z⁴S° c⁴D	0.646 01	0.6075	3d ⁶ ³ F4d	fD	0.232 40*	0.2007
		0.639 58	0.6191	a3d ^{6 3} F4d		0.231 06*	0.1940
3d ^{6 3} P4p	y⁴P°	0.635 49	0.5997		⁴G ⁴G	0.231 07	0.1946
3d ^{6 3} F4p	y ⁴ F°	0.623 42	0.5947	3d ^{6 5} D5d		0.230 88	0.2389
3d ^{6 3} H4p	z ⁴ G°	0.635 48	0.6168	3d ⁶ ³ Fa4d	⁴H	0.228 98	0.1939
3d ^{6 3} H4p	z ⁴ H°	0.634 34	0.6193	3d ⁶ ³ F4d	⁴F	0.216 64	0.1875
3d ⁶ ³ H4p	z ⁴ I°	0.629 42	0.6194	3d ⁶ ⁵ D5d	⁴p	0.215 10	0.2267
3d ⁶ ³ P4p	y⁴D°	0.621 32	0.5896	3d ⁵ 4s ³ D4p	⁴D°	0.213 54	0.2129
3d ^{6 3} F4p	x ⁴ D°	0.613 70	0.5857	3d ⁵ 4s ³ D4p	⁴P°	0.219 16*	0.2129
3d ^{6 3} F4p	y " G"	0.606 78	0.5844	3d ⁶ ³ F4d	⁴F	0.21661	0.1875

 Table 3. (continued)

			E (Ry)			E ((Ry)
State		Obs.	Calc.	State		Obs.	Calc.
3d ⁶ ³ G4p	x⁴G°	0.590 33	0.5723	3d ^{6 5} D6p	⁴ D°	0.213 08*	0.2101
3d ⁶ ³ G4p	x ⁴ F°	0.585 42	0.5645	3d ^{6 5} D6p	۴F°	0.212 78	0.2133
3d ⁶ ³ G4p	y ⁴ H°	0.583 58	0.5669	3d ^{6 5} D6p	⁴P°	0.209 37	0.2101
3d5 4s5S4p	x ⁴ P°	0.558 89	0,5729	3d ⁵ 4s ³ F4p	⁴G°	0.208 68*	0.2115
3d ⁶ ³ D4p	w ⁴ P°	0.533 29	0.5009	3d ⁵ 4s ³ F4p	⁴D°	0.202 48*	0.1984
3d ^{6 3} D4p	w ⁴ F°	0.529 87	0.5007	3d ^{6 3} P5p	4S°	0,202 14	0.1919
3d ⁶ ³ D4p	w⁴D°	0.528 17	0.4962	3d ⁶ ³ G4d	'G	0.201 97	0.1830
3d54s2	⁴F	0.520 31	0.4884	3d6 3G4d	⁴H	0.201 81	0.1830
3d ^{6 5} D5s	e ⁴ D	0.462 40	0.4555	3d ⁵ 4s ³ F4p	⁴F°	0.201 41	0.2022
3d ^{6 5} D4d	f [‡] D	0.416 38	0.4225	3d54s3G4p	⁴G°	0.200 38	0.1948
3d ^{6 5} D4d	e⁴G	0.413 10	0.4195	3d6 3G4d	4]	0.200 19	0.1812
3d ^{6 5} D4d	4S	0.408 48	0.4142	3d ⁶ ³ G4d	٠ D	0.200 03	0.1808
3d ^{6 5} D4d	e ⁴ F	0.402 42	0.3933	3d ⁶ ³ D5s	⁴D	0.198 84	0.1860
3d ⁶ ³ P4p	v ⁴ D°	0.399 18	0.3401	3d ³ 64s ³ G4p	4H°	0.198 10	0.2010
3d ^{6 5} D4d	⁴P	0.387 09	0.3747	3d ⁶ ³ P5p	4p0	0.195 11	0.1868
b3d ^{6 3} F4p	⁴Ĝ°	0.369 44	0.3475	3d ⁶ ³ H5p	4I°	0.190 04	0.1751
3d ^{6 3} P4p	4S°	0.363 82	0.3144	3d ⁶ ³ H5p	⁴G°	0.188 33	0.1731
3d ^{6 5} D5p	⁴D°	0.363 36	0.3631	3d ⁶ ³ P5p	⁴D°	0.187 67*	0.1802
3d ^{6 5} D5p	⁴F°	0.362 61	0.3654	3d ⁶ ³ G4d	⁴F	0.185 85	0.1695
3d D5p 3d6 5D5p	⁴ p °	0.355 18	0.3575	3d ^{6 3} F5p	⁴F°	0.185 29	0.1664
30° D5p	⁴p°	0.355 18		3d ⁵ 4s ³ G4p	r ⁴H°	0.183 29	0.1380
3d ⁶ ³ P4p			0.3348	30°48°G4p	⁴D°		
3d54s5G4p	x⁴H°	0.349 80	0.3693	3d ⁶ ⁴ F5p		0.182 37*	0.1537
3d ⁵ 4s ⁵ G4p	v ⁴ F°	0.348 10	0.3618	3d ⁶ ³ H5p	⁴ H°	0.172 27	0.1821
3d ^{6 3} F4p	⁴D°	0.343 16	0.3100	3d ⁶ ³ F5p	⁴G°	0.169 26	0.1498
3d ^{6 3} F4p	u ⁴ F°	0.338 17	0.2981	3d ⁵ 4s ³ H4p	4I°	0.167 56*	0.1444
3d ⁵ 4s ⁵ G4p	w⁴G⁰	0.332 03	0.3085	3d ⁶ ³ G5p	⁴F°	0.166 97	0.1575
3d ⁵ 4s ⁵ P4p	⁴ p °	0.328 56	0.2877	3d ³ 4s ⁵ F4p	⁴G°	0.165 94*	0.1777
3d ⁵ 4s ⁵ P4p	⁴D°	0.314 57	0.2952	3d ^{6 5} D7s	⁴D	0.158 33	0.1614
3d ⁶ ³ P5s	⁴P 4	0.296 95	0.2832	3d ⁶ ⁵ D6d	⁴F	0.149 40*	0.1396
3d ^{6 3} H5s	e⁴H 4=2	0.293 68	0.2404	3d ⁶ ³ G5p	⁴H°	0.148 92*	0.1455
3d ⁵ 4s ⁵ P4p	4S°	0.293 58	0.2718	3d ⁶ ⁵ D6d	⁴G ⁴~~	0.146 62	0.1538
3d ⁵ 4s ⁵ D4p	4F°	0.292 37	0.2822	3d ⁶ ³ G5p	⁴Gº	0.142 92*	0.1402
3d ⁵ 4s ⁵ D4p	⁴D°	0.291 53	0.2782	3d ⁶ ³ D4d	⁴F	0.141 46*	0.1158
3d ^{6 3} F5s	f ⁴ F	0.218 08	0.2401	3d ⁵ 4p ²	⁴P	0.11286*	0.1231
3d⁵4s⁵D4p	⁴P°	0.263 75	0.2558				
			Double	ets: 62-cc			
3d ⁷	a ² G	1.043 19	0.9380	3d6b3P4p	u^2D^o	0.346 74	0.2851
3d ⁷	a ² P	1.020 79	0.9148	3d ⁶ b ³ F4p	$^{2}D^{\circ}$	0.326 72*	0.2698
3d ⁷	a ² H	1.002 42	0.9222	3d ⁶ b ³ P4p	² P°	0.323 64*	0.2902
3d ⁷	a ² D	0.999 85	0.8896	3d ⁶ b ³ F4p	²F°	0.323 40	0.3180
3d ⁶ a ³ P4s	b ² P	0.951 23	0.8763	3d ⁵ 4s ³ G4p	²H°	0.313 58	0.3181
3d ⁶ ³ H4s	b ² H	0.950 46	0.8958	3d ⁵ 4s ³ G4p	²F°	0.311 93	0.2582
3d ⁶ a ³ F4s	a ² F	0.939 60	0.8905	3d ⁵ 4s ³ P4p	²p°	0.302 80*	0.2533
3d ⁶ ³ G4s	b²G	0.911 26	0.8667	3d°b¹G4p	²H°	0.295 89	0.2394
3d ⁷	b ² F	0.898 83	0.7842	3d°b'G4p	²F°	0.288 47*	0.2350
3d ⁶ 114s	a ² I	0.889 97	0.8470	3d ⁶ ³ H5s	e²H	0.285 71	0.2324
3d ⁶ a ¹ G4s	c²G	0.884 59	0.8307	3d ⁶ ³ P5s	² P	0.284 33	0.2750
3d a O4s	b ² D	0.859 80	0.7966	3d54s3G4p	²Ġ°	0.281 66	0.2741
3d ⁶ a ¹ S4s	a ² S	0.850 46	0.7887	3d*b'G4p	²G°	0.280 44	0.2090
3d ⁶ a ¹ D4s	c ² d	0.841 74	0.7565	3d 5 4s 3P4p	$^{2}D^{\circ}$	0.274 78*	0,2521
3d a D4s 3d ^{6 1} F4s	c ² F	0.780 35	0.6907	3d ⁶ a ³ F5s	e ² F	0.272 94	0,2314
3d ⁷ 74s	¢ F d²D	0.753 26	0.6128	3d 4s 3D4p	$^{2}D^{\circ}$	0.255 92	0.2379
3d ⁶ b ³ P4s	c ² P	0.733 20	0.5975	3d 4s D4p 3d 54s 3D4p	²F°	0.233 92	0.2101
	d ² F		0.5997	3d 4s D4p 3d ⁶ ³ G5s	e ² G	0.244 03	0.2310
3d ⁶ b ³ F4s	ur	0.689 51	Ų. J. J. J. I	λα (1)29	τ U	V.674 UJ	0.6310

Table 3. (continued)

		E	(Ry)			E (Ry)
State		Obs.	Calc.	State		Obs.	Calc.
3d6b1G4s	d²G	0.655 27	0.5513	3d ⁵ 4s ³ P4p	2S°	0.242 28	0.2123
3d ⁶ a ³ P4p	z^2D°	0.629 21	0.5944	3d ⁶ ³ H4d	2K	0.240 06	0.2092
3d ⁶ ³ H4p	z^2G°	0.622 99	0.6072	3d ⁶ ³ H4d	²F	0.238 66	0.2066
3d ⁶ ³ H4p	$z^2 l^{\circ}$	0.620 49	0.5575	3d ^{6 3} H4d	²G	0.233 35	0.2008
3d ⁶ a ³ F4p	z^2F°	0.603 34	0.5794	3d ^{6 3} H4d	²[0.235 71	0.2061
3d ⁶ a ³ P4p	z^2P^o	0.598 97	0.5633	3d ^{6 3} P4d	² p	0.231 74	0.2095
3d ⁶ a ³ F4p	y ² G°	0.597 78	0.5713	3d ⁶ ³ P4d	2 D	0.230 69*	0.2108
3d ⁶ ³ H4p	z²H°	0.593 26	0.5772	3d ⁶ a ³ F4d	²H	0.224 86	0.1917
3d ⁶ a ³ P4p	z^2S°	0.586 00	0.5491	3d ^{6 3} H4d	² H	0.223 13	0,1896
3d ⁶ a ³ F4p	y^2D°	0.578 15	0.5498	3d ⁶ a ³ F4d	²F	0.221 86*	0.1920
3d ⁶ ³ G4p	y ² H°	0.572 44	0.5581	3d ⁵ 4s ³ D4p	² P°	0.220 47	0.2001
3d ⁶ ³ G4p	y ² F°	0.555 23	0.5278	3d ⁶ 4s ³ I4p	²K°	0.220 43	0.2409
3d ⁶ ³ G4p	x²G°	0.548 10	0.5262	3d ⁶ 4s ³ I4p	² H°	0.216 16	0.2158
3d ⁶ ¹ I4p	z²K°	0.540 65	0.5211	$3d^{6}a^{3}F4d$	² G	0.215 43	0.1849
3d ⁶ a ¹ G4p	x2H°	0.531 75	0.5012	$3d^6a^3F4d$	²D	0.21085	0.1754
3d ⁶ a ¹ G4p	x²F°	0.524 18	0.4907	3d ⁶ a ³ F4d	² P	0.204 57*	0.1648
3d ⁶ a G4p	w^2G^o	0.523 43	0.4883	3d ^{6 1} I5s	²Į	0.199 70	0.1374
3d ⁶ 3D4p	y ² P°	0.522 76	0.4852	3d ^{6 3} P5p	² D°	0.197 03*	0.1840
3d ⁶ ¹ I4p	w ² H°	0.518 36	0.4880	3d ⁶ ³ G4d	2Į	0.194 54	0.1786
3d ⁶ ¹ I4p	y²ľ°	0.515 65	0.4927	3d ⁵ 4s ³ 14p	²ĭ°	0.194 54	0.2054
3d ⁶ 3D4p	x2D°	0.510 23	0.4747	3d ⁶ ³ D5s	2D	0.190 66	0.1316
3d ⁶ ³ D4p	w^2F^o	0.499 55	0.4606	3d ^{6 3} G4d	² G	0.189 60	0.1638
3d ⁶ a S4p	x ² P°	0.494 60	0.4429	3d ^{6 3} G4d	² H	0.188 85	0.1625
3d ⁵ 4s ²	²H	0.485 92*	0.4588	3d ^{6 3} H5p	2G°	0.185 30	0.1452
3d ⁶ a ¹ D4p	v^2F°	0.479 20	0.4212	3d ^{6 3} G4d	^{2}D	0.185 26	0.1646
3d54s2	²G	0.475 24	0.4459	3d ⁶ ³ H5p	² I ^o	0.186 64	0.1231
3d ⁶ a ¹ D4p	W^2D^0	0.473 36	0.4134	3d ⁶ ³ G4d	²F	0.181 95	0.1519
3d ⁶ a 1D4p	w^2P^o	0.468 80	0.4130	3d ⁵ 4s ³ F4p	² G°	0.181 28*	0.1823
3d ⁵ 4s ²	² F	0.445 25	0.4881	3d ^{6 3} H5p	² H°	0.180 07	0.1370
3d ^{6 1} F4p	v²G°	0.427 70	0.3710	3d ⁶ ³ F5p	²G°	0.172 93	0.1399
3d ⁶ ¹ F4p	v^2D^o	0.423 64	0.3630	3d ^{6 3} F5p	²F°	0.171 92	0.1417
3d ⁶ F4p	u2F°	0.401 36	0.3411	3d ⁶ 3G5p	²H°	0.145 10	0.1415
3d°b¹P4p	2S°	0.378 64	0.3210	3d ⁶ ³ G5p	2G°	0.142 85*	0.1310
3d ⁶ b ³ F4p	п ₅ С ₀	0.348 02	0.2886	•			

The comparison needs to be viewed in light of the following: while the present calculations do not explicitly account for the relativistic effects, some allowance is made by using observed energies to obtain the f-values with improved accuracy. The electron correlation effects should be better represented in the present work, based on *ab initio* close coupling calculations, than the semi-empirical method of Kurucz involving fitting of parameters to observed energy levels. For most transitions therefore the present data for Fe II obtained in this manner should be at least as accurate as currently available values. The present results provide an alternative dataset for a large number of oscillator strengths for Fe II, although it is difficult to state the uncertainties precisely. The LS coupling f-values given in table 4(a) provide an overall indication of the accuracy of the total multiplet strengths in the total dataset,

In table 4(b) we further extend the comparisons to the fine structure components, obtained through algebraic transformation (Alien 1976) as described earlier. Only selected oscillator strengths are presented for which a complete or almost complete set of fine structure oscillator strengths is available.

Table 4. (a) Comparison of oscillator strengths, f_{if} , of Fe II in LS coupling. ΔE is the transition energy, (b) Comparison of fine structure oscillator strengths for transitions in Fe II. For each transition the first line corresponds to the LS values and the following lines to the fine structure components, (c) Calculated and measured lifetimes, τ , of Fe II levels.

	ΔE		$f_{d'}$			ΔE		Su .
Transition	(Ry)	Present	NIST	Kurucz	Transition	(Ry)	Present	Kurucz
a °S →z °P°	0.180	0.0434	0.041	0.0196	a °D →z °P°	0.389	0.126	0.094
a °D →z °D°	0.350	0.2945	0.26	0.297	a °D →z °F°	0.382	0.399	0.413
a⁴P →z⁴P°	0.306	0.0838	0.066	0.106	z ⁶ P° →e ⁶ D	0.321	0.142	0 149
a ¹P →z ⁴D°	0.284	0.0273	0.0168	0.0289	$b^4P \rightarrow y^4P^0$	0.359	0.210	0.263
b ⁴P →z⁴D°	0.213	8.0 (-5)	0.0028	0.0029	a⁴D →z⁴D°	0.332	0.281	0.273
a⁴D →z⁴P°	0.354	0.142	0.13	0.152	b⁴D →z⁴P°	0.144	1.12 (-3)	1.10(-3)
a⁴D →z⁴F°	0.332	0.378	0.33	0.351	b⁴D →z⁴D°	0.121	4.2 (-4)	4.5 (-4)
a⁴F →z⁴F°	0.385	0.0431	160.0	0.050	a ⁴F →z ⁴D°	0.386	0.085	0.0804
b⁴F →z⁴F°	0.200	8.7 (-4)	7.1(-4)	8.6 (-4)	a ⁴ H →z ⁴ H°	0.360	0.243	0.255
$a^2S \rightarrow z^2P^\circ$	0.251	0.0132	0.0084	0.023	a ⁴H →z ⁴G°	0,359	0.170	0.132
a ² S → v ² P°	0.328	0.112	0.0087	0.265	b ⁴ P → z ⁴ S°	0,348	0.0663	0.0087
a²S →x²P°	0.356	0.535	0.39	0.536	$b^2D \rightarrow y^2P^0$	0.337	0.120	0.108
² P →z ² P°	0.352	0.208	0.23	0.26	b²D →x²D°	0.350	0.276	0.261
$\begin{array}{ccc} & P & \rightarrow z^2 P^{\circ} \\ & P & \rightarrow x^2 P^{\circ} \\ & P & \rightarrow x^2 P^{\circ} \end{array}$	0.347	0.0557	0.098	0.035	a ²F →y ²G°	0.342	0.337	0.280
$z^2F \rightarrow z^2F^0$	0.336	0.202	0.14	0.109	$c^2F \rightarrow v^2D^n$	0.357	0.182	0.168
b²F →x²G°	0.351	0.0661	0.042	0.0668	d²F →u²G°	0.341	0.223	0.183
°F →v °G°	0.353	0.342	0.31	0.349	b²P →z²D°	0.322	0.106	0.217
² G →y ² F°	0.356	0.182	0.13	0.147	$b^2P \rightarrow z^2S^n$	0.365	0.0036	0.080
o G → v ² G°	0.313	1.0(-4)	3.8 (-3)	3.8 (-3)	$b^2D \rightarrow z^2D^o$	0.231	1.3(-3)	1.4(-3)
G →y²H°	0.339	0,230	0.18	0.21	b²H →z²H°	0.357	0.161	0.0946
² G →z ² F°	0.281	9.1 (-3)	5.2 (-3)	6.0(-3)	b²H →z²I°	0.330	0.158	0.278
2G →x2H°	0.353	0.140	0.17	0.278	a ² l →z ² H°	0.297	0.119	0.110
b²H →z²G°	0.327	0.0122	0.069	0.108	c ² G →z ² G°	0.262	6.5 (-3)	6.7(-3)

(b)

	λ_{if}				Su		la har				f_{tf}	
Transition	(Å)	g_i	8/	Present	NIST	Kurucz	(Å)	gı	8/	Present	NIST	Kurucz
a ⁶ S° → z ⁶ P°		6	18	0.0434	0.041	0.0196						
	5170	6	8	9810.0	0.023	0.0083	4925	6	4	0.0099	0.008	0.0046
	5020	6	6	0.0146	010.0	0.0066						
$a ^6D^c \rightarrow z ^6P^o$		30	18	0.126		0.0938						
	2344	10	8	0.126	11.0	0.129	2360	4	б	0.0377		0.0718
	2366	8	8	0.0449	0.051	0.0566	2328	6	4	0.0396	0.032	0.039
	2381	6	8	0.0099	0.036	0.0392	2339	4	4	0.0887	0.087	0.103
	2334	8	6	0.0814		0.0805	2345	2	4	0.126	0.13	0.177
	2349	6	6	0.077		0.094						
a °D° →z °D°	,	30	30	0.294	0.26	0.297						
	2600	10	10	0.240	0.22	0.261	2608	6	4	0.118	0.11	0.133
_	2587	10	8	0.0548	0.065	0.0766	2632	4	6	0.175	0.12	0.133
	2626	8	10	0.0675	0.043	0.0459	2621	4	4	0.00322	0.0037	0.003 95
	2613	8	8	0.131	0.11	0.135	2615	4	2	0.114	0.10	0.119
	2599	8	6	0.0948	0.099	0.117	2629	2	4	0.227	0.18	0.188
	2632	6	8	0.125	0.084	0.0887	2622	2	2	0.0652	0.050	0.0587
	2618	6	6	0.0502	0.045	0.0517						
a D° →z°F°		30	42	0.397		0.413						
	2383	10	12	0.342	0.328	0.36	2400	6	6	0.131	0.12	0.130
	2374	10	10	0.0527	0.028	0.0340	2396	6	4	0.0227	0.019	0.017
	2368	10	8	0.004 25		4.4(-5)	2411	4	6	0.203	0.19	0.232
	2396	8	10	0.289	0.27	0.323	2407	4	4	0.161	0.14	0.167
	2389	8	8	0.0968	0.089	0.0924	2405	4	2	0.0314	0.031	0.0290
	2384	8	6	0.0122		0.0057	2414	2	4	0.175	0.19	0.200
	2406	6	8	0.243	0.20	0.265	2412	2	2	0.220	0.21	0.236
$a^4P^c \rightarrow z^4P^c$		12	12	0.0838	0.066	0.106						
	2986	6	6	0.0585	0.048	0.0763	2945	4	2	0.0353	0.03	0.0452
	2949	6	4	0.0254	0.017	0.0327	2986	2	4	0.0696	0.048	0.0843
•	3004	4	6	0.0374	0.029	0.0427	2965	2	2	0.0140	0.012	0.0169
	2966	4	4	0.0112	0.0079	0.0154	_, ,,,	_	_			
$a^4P^c \rightarrow z^4D^\circ$	2,00	12	20	0.0273	0.0168	0.0289						
	3229	6	8	0.0217	0.012	0.023	3188	4	4	18800.0	0.0049	0.008 34
	3194	6	6	0.00495	0.0019	0.003 36	3171	4	2	0.00139	6.9(-4)	0.001 16

 Table 4. (continued)

	λ,,		-		f_{ψ}		λ,,				f_{ij}	
Transition	(Å)	8 i	g f	Present	NIST	Kurucz	(Å)	g,	81	Present	NIST	Kurucz
	3168	6	4	5.5 (-4)	1.4 (-4)	2.8 (-4)	3211	2	4	0.0137	1800.0	0.0170
	3214	4	6	0.0172	0.015	0.0206	3195	2	2	0.0137	0.0095	0.0153
a ⁴D° →z ⁴P°		20	12	0.142	0.13	0.152						
	2563	8	6	0.143	0.11	0.144	2583	4	4	0.0754	0.077	0.0885
	2592	6	6	0.0423	0.052	0.055	2595	2	4	0.0235	0.026	0.0306
	2612	4	6	0.0070	0.0093	110.0	2568	4	2	0.0593	0.056	0.0604
	2564	6	4	0.0997	0.085	0.0995	2579	2	2	0.118	0.13	0.132
a ⁴ D° →z ⁴ D°		20	20	0.281		0.273						
	2740	8	8	0.241	0.22	0.268	2770	4	6	0.0973	0.0077	0.0267
	2715	8	6	0.0404	0.045	0.05	2750	4	4	0.112	0.12	0.131
	2774	6	8	0.0528	8.0 (-5)	0.0049	2738	4	2	0.0703		0.0713
	2748	6	6	0.161	0.18	0.189	2763	2	4	0.139	0.026	0.0632
	2728	б	4	0.0659	0.063	0.0746	2750	2	2	0.140	0.12	0.131
a ⁴ D" →z ⁴ F°		20	28	0.378	0.33	0.351		_	-			*****
	2757	8	10	0.336	0.30	0.266	2710	6	4	0.00365	1.9 (~4)	2.3 (-4)
	2718	8	8	0.0389	0.0011	0.0041	2747	å.	6	0.303	0.33	0.364
	2694	ž	6	0,00199	100.0	7.2 (-5)	2732	4	4	0.0760	0.028	0.0485
	2750	6	8	0.308	0.32	0.351	2744	2	4	0.378	0.41	0.446
	2726	6	6	0.0662	110.0	0.0264	2177	-	•	0.575	0,41	0,440
b ⁴ D° →x ⁴ D°	2,20	20	20	0.0223	0.011	0.02						
0 0 7 0	3178	8	8	0.0189	0.012	0.017	3134	4	6	0.00782	0.0035	0.0037
	3146	8	6	0.00317	0.0023	0.0032	3115	4	4	0.00899	0.0039	0.0099
	3169	6	8	0.00377	0.0023	5.8 (-4)	3106	4	2	0.00564	0.0057	0.0058
	3136	6	6	0.0042	0.012	0.014	3116	2	4	0.0112	0.0037	0.0038
	3117	6	4	0.00524	0.0068	0.0058	3106	2	2	0.0112	0.011	8010.0
a ⁴F° →z ⁴D°	3117	28	20	0.00524	0.0008	0.0804	2100	4	2	0.0113	0.011	0.0108
ar -42 D	2349	íô	8	0.0859	0.034	0.0304	2400	4	6	0.00168		0.0041
	2380	8	8		0.034	0.0754	2369	6	4	0.00168	0.023	0.0609
	2403	6	8	0.0121	0.13	0.0237		-	4		0.033	
	2361		6	8.1 (-4)	0.0022	0.066	2385 2376	4	2	0.0237	0.020	0.036
	2384	8 6	6	0.0733 0.0206			2370	4	2	0.0595	0.041	0.0648
a 'F° →z 'F°	2384	28	28		0.029 <u></u> 0.031	0.035						
ar →zr	2361	10		0.0431	0.031	0.030	2386	,		0.00540	0.0000	0.006.60
	2332		10 8	0.0396		0.0429		6	8	0.00763	0.0038	0.006 57
	2332	10	-	0.00362	0.019		2367	6	6	0.0296	0.0084	0.0191
	2363	8	10	0.00442	0.0029	0.00504	2356	6	4	0.00577	0.013	0.0150
		8	8	0,0329	0.011	0.0227	2383	4	6	0.00856	0.0051	0.007 14
. 2mt _ 2mn	2345	8	6	0.00583	0.018	0.0208	2371	4	4	0.0344	0.012	0.0265
a²S° →z²P°	0.000	2	6	0.0132	0.0084	0.023	2/2/			0.00404	0.000=	
200 200	3622	2	4	0.00878	1200.0	0.015	3626	2	2	0.00438	0.0035	0.008
a²S° →y²P°	~~~		6	0.113	0.0087	0.265		_	_			
. Zoe 2-0	2781	2	4	0.0749	0.054	0.085	2781	2	2	0.0374	0.034	0.18
a²S° →x²P°	0.575	2	6	0.535	0.39_	0.536		_	_			
	2571	2	4	0.355	0.23	0.347	2541	2	2	081.0	0.15	0.188

(c)

			τ (ns)					τ (ns)	
State	j	Present	Expt	Kurucz	State	J	Present	Expt	Kurucz
z ⁶ P°	7/2	3.884	3.73(0.06)*, 3.8(0.2)* 3.5(0.3)*, 3.73(0.05)*	3.28	z °F°	9/2	3.024	3.2(0.2) ^b , 3.24(6) ^d	2.89
r ⁶ P°	5/2	3.844	3.79(0.12), 3.7(0.2)	3.26	z°F°	7/2 5/2	3.034 2.973	3.26 (10) ^d 3.3(0.2) ^e 3.33(9) ^d	2.92 2.93
z ⁶ P°	3/2	3.819	3.5(0.3)°, 3.83(0.07)° 3.71(0.12)°, 3.6(0.2)°	3.25	z °F° ≥ °F°	3/2 1/2	2.908 2.878	3.3(0.2)°, 3.34(10)° 3.3(0.3)°	2.94 2.94
4D°	9/2	3.460	3.4(0.3) ^e 3.7(0.2) ^b , 3.7(0.2) ^e	3_41	z⁴P° z⁴D°	5/2 7/2	3.250 2.476	3.44(0.11)* 3.02(0.07)*, 3.1(0.2)b	2.43
ʻD°	7/2	3.487	3.75(20) ⁶ , 3.8(0.3) ⁶ 3.68(0.7) ^d	3,43	z⁴D° z⁴D°	5/2 3/2	2.496 2.494	3.1(0.08)*, 3.1(0.2)* 3.0(0.2)*	2.44 2.43
z °D°	5/2 3/2	3.406 3.392	3.7(0.2) ⁶ , 3.63(8) ^d 3.7(0.2) ⁶ , 3.7(0.2) ^c	3.44 3.45	z ⁴D° z ⁴F°	1/2 9/2	2.498 3.471	2.9(0.2) ^b 3.87(0.09) ^a , 3.7(0.2) ^b	2.42 3.34
z ⁶ D°	1/2	3.259	3.83(10) ^d 3.8(0.2) ^b , 3.8(0.3) ^c	3,45	z fr° z fr°	7/2	3.435 3.417	3.63(0.11)*, 3.6(0.2)*	3.22
			3.76(10) ⁶		z F°	5/2 3/2	3.422	3.75(0.14)*, 3.7(0.2)* 3.7(0.2)*	3.26 3.3
, ⁴F°	11/2	2.982	3.2(0.2) ⁶ , 3.3(0.2) ^c 3.19(4) ^d	2.83					

Guo et al (1992). Hannaford et al (1992).

^c Schade et al (1988). ^d Biemont et al (1991).

There are a few measured values of Fe II that are with low uncertainty, rated B or <10%. However, most of them are rated with a 50% uncertainty (D in the NIST compilation). The measured R-values of the fine structure transitions of an LS multiplet usually do not correspond to the same experiment, rather to more than one experimental source. Among all the transitions, one complete set that has been rated with 10% uncertainty is a $^6D^e \rightarrow z$ $^6D^o$. For this particular multiplet, the component fine structure transitions show about same level of agreement between the present values and the measured ones, as between the Kurucz values and the experimental values. However, overall comparison shows that the present values agree better with the measured values than the Kurucz data set. All the transitions listed in the NIST compilation have been compared, and this provides more detailed information on the uncertainties than the comparison of total LS multiplets. (The wavelengths given in table 4(b) are approximate.)

Another interesting comparison may be made between the calculated and observed lifetimes. We calculate the lifetime of a state based on all allowed transitions from the given state, i.e., $\tau_j = [\Sigma A_{Ji}]^{-1}$ where $A_{ji} = a^3 g_{i}/g_{j} E_{ji}^2 f_{ij}/\tau_0$ is the transition probability in s^{-1} , f_{ij} is the oscillator strength, τ_0 is the unit of time, and the sum is over all lower i-states that are connected by dipole allowed transitions from the upper state j. The lifetimes are calculated using the present fine structure f-values. Table 4(c) presents the calculated lifetimes and compares with the available measured values, and those from Kurucz. It should be noted that the Kurucz calculations involve forbidden transitions, and hence the lifetimes include both the dipole allowed and forbidden transitions; however the latter contribution is expected to be small for most levels under consideration. Present lifetimes of the fine structure levels of z $^6P^0$ agree well with the recent measured values of Hannaford et at (3992) and Guo et at (1992), whereas those of Schade et at (1988) are lower than the present values and other measured values. Values obtained by Kurucz are also lower than the present ones in a similar manner. For other states, the agreement between the present values and the experimental ones is better than that of Kurucz. The present values are in general within 5-10% of the experimental values.

4.3. Photoionization cross sections

Photoionization cross sections, including the detailed autoionizing resonances, are calculated for all bound states which include the 743 bound states of Fe II that lie below the first ionization threshold. Selected examples of the cross sections are presented. Figure 1 shows the photoionization cross sections of the 3d⁵4s ⁶D ground state of Fe II. As discussed by LeDourneuf *et al* (1993) the present ground state cross sections are about two orders of magnitude higher than the earlier calculations (Sawey and Berrington 1992) which included only the ground state of Fe III in the target expansion, whereas the dominant contribution stems from the photoionization of the 3d shell i.e. through coupling to the excited 3d⁵4s state of Fe III.

Figure 2(a) shows the photoionization cross sections of the lowest three quartet states, $3d^7(^4F)$, $3d^64s(^4D)$ and $3d^7(^4P)$. These metastable states are likely to be of considerable importance in astrophysical plasmas as one expects these to be in local thermodynamic equilibrium (LTE) even at fairly low densities owing to small radiative decay rates. Thus there may be significant populations in the low-lying metastable states,

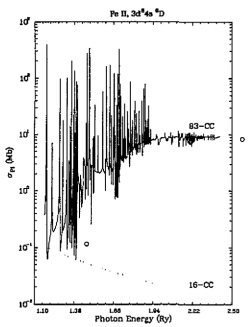


Figure 1. Photoionization cross section of the 3d⁵4s⁶D ground state of Fe II using 83-CC expansion. The dotted curve corresponds to earlier 16-CC calculations (Sawey and Berrington 1992); also shown are the central field cross sections in circles (Reilman and Manson 1979).

comparable to the ground state, in accordance with respective statistical weights. *Photoionization models therefore need to consider not only ground state photoionization but also that of at least these three states* (other states lie significantly higher and would be relatively less important). Figure 2(b) presents the photoionization cross sections of the three lowest doublet states of Fe II. Extensive resonances can be seen to dominate the metastable states $3d^7$ P and $3d^7$ P for the entire energy range up to the highest target threshold, and resulting in large variations in the background cross sections. Though the background cross sections smooth out at higher energies due to weaker coupling to high target states, all $3d^7$ states and 3d64s D show significant variations in the cross sections in the near threshold region due to resonances.

Figure 3 presents photoionization cross sections of two octet states, 3d⁵4s ⁷S5s ⁸S and the equivalent electron state 3d⁵4p² ⁸P. Both states show the presence of a large resonance at the threshold, and the repetition of resonance patterns converging on to the ⁷P^o target state. The octet state photoionization cross sections have been obtained for the first time.

Figure 4 consists of eight panels presenting photoionization cross sections of the Rydberg series of states, $3d^6$ ⁵Dnp ⁴ P^o where $4 \le n \le 11$ of Fe II. The figure illustrates the wider photo-excitation of core (PEC) resonances which occur at the energies of the target states $3d^5$ ⁶S4p (⁵P°), $3d^5$ ⁴P4p(⁵P°), $3d^5$ ⁴P4p(⁵P°), $3d^5$ ⁴F4p(⁵D°), corresponding to dipole allowed transitions from the ground state ⁵D (the positions of the PEC resonances are pointed out by arrows in the top panel). At these energies, the outer 'np'electron remains as a spectator while the ground core state $3d^54s$ ⁵D is excited via strong dipole transitions giving rise to wide resonances in photoionization cross sections (Yu and Seaton 1987).

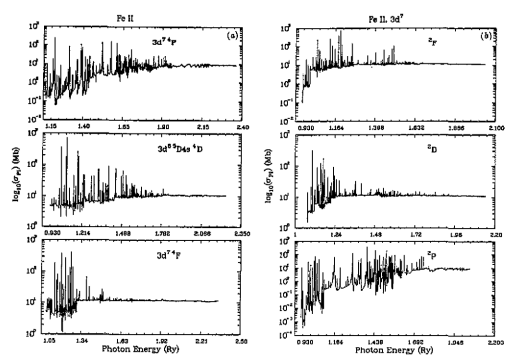


Figure 2. Photoionization cross sections of the metastable states of Fe II: (a) first three lowest quartet states, (b) three lowest doublet states.

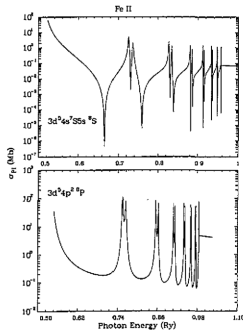


Figure 3. Photoionization cross sections of the two octet states, ⁸S and ⁸P, of Fe II.

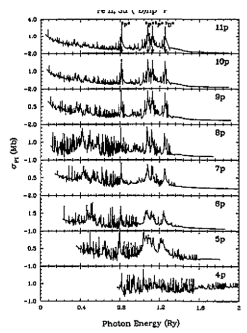


Figure 4. Photoionization cross sections of bound states along a Rydberg series, $3d6(^5D)np^4P^o$, with $4 \le n \le 11$ illustrating photoexcitation-of-core (PEC) resonances.

These resonances appear to be better resolved at higher excited states since the Rydberg resonances are usually weaker. The phenomenon of PEC resonances contradicts the usual assumptions of smooth hydrogenic behaviour of excited state photoionization cross sections, which may in fact be quite non-hydrogenic owing to the PEC features. It should be noted that even though there are more quintet states of Fe III in table 1 that are accessible through dipole transitions from the ground state, the figure shows only four since we treat several excited states to be degenerate as explained in section 3 and specified in the target states column of table 1.

4.4. Monochromatic opacities

The primary goal of the Opacity project is to calculate stellar opacities. Thus it is of some interest to examine the contribution to the opacities of a large calculation such as for Fe II reported herein. The total monochromatic opacity K_V is obtained on summing the contributions from all of the radiative processes such as bound-bound transitions, bound-free transitions, Thomson scattering, and free-free transitions (the former two are the main contributors). The theoretical steps involved in the opacities calculations are summarized by Seaton (1987). The contribution to the monochromatic opacity from a bound-bound transition is obtained as

$$\kappa_v = 2\pi^2 e^2/(mc) N_i f_{ii} \phi_v$$

where N_l , is the number density of the ion, and Φ_v is a profile factor normalized to $\int \Phi_v dv = 1$; and that of a bound-free transition is

$$\kappa_{\nu} = N_{\nu} \sigma_{\rm PI}$$

Where O_{PI} is the photoionization cross section. In terms of the monochromatic opacities, the flow

of radiation through a plasma is governed by the Rosseland mean defined as

$$\frac{1}{\kappa_{R}} = \int_{0}^{\infty} \frac{1}{\kappa_{\nu}} g(u) \, \mathrm{d}u \tag{1}$$

where

$$u = \frac{hv}{kT}$$
 $g(u) = \frac{15}{4\pi^4}u^4 \frac{\exp(-u)}{[1 - \exp(-u)]^3}$.

The Rosseland mean is the weighted harmonic mean of the monochromatic opacities and K_R is the Rosseland mean opacity. We have carried out separate calculations for the monochromatic and the mean opacities (using the OP code OPAC by Y Yu), for the single ion Fe II, with the new *R*-matrix data for the bound-bound and bound-free transitions from this work, and with the earlier *R* -matrix data of Sawey and Berrington (1992). At a temperature of 16 000 K and a density of 10^{16} cm⁻³, we obtain values of K_R to be 185 and 120 respectively, with the new and old data, resulting in an increase of over 50%. The detailed monochromatic opacity spectrum of Fe II, with the two data

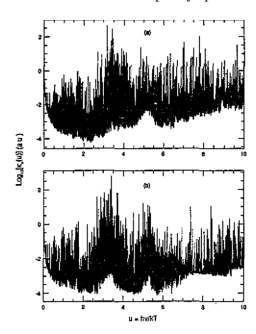


Figure 5. Monochromatic opacities, K_v , of Fe II at $\log T = 4.2$ and $\log N_e = 16$, (a) using the present radiative data and (b) the earlier data (Sawey and Berrington 1992).

sets, is shown in figure 5. In particular we note the rather large gap in the spectrum in the earlier data (bottom panel) around u = 3.5 (5 eV), which has been filled in to a large extent by the new Fe II data (top panel). The Rosseland mean opacity, being a harmonic mean over the monochromatic opacities, is very sensitive to gaps or holes in the detailed spectrum, at low values of hv/kT (equation (1)), and the feature in figure 5 is a significant contributor to the enhanced opacity. Also, in the high energy region, present opacities exhibit a rising trend and are more enhanced than those obtained using the earlier data.

4.5. Conclusion

Extensive radiative calculations for Fe II are described and it is expected that the new data will be applicable to a variety of astrophysical applications, in addition to the original aim of the calculations of accurate plasma opacities. The *LS* multiplet and fine structure oscillator strengths are compared with the available data and shown to be generally more accurate than other previous theoretical calculations (although there may be significant uncertainties for the weaker transitions). Most of the detailed photoionization data have been calculated for the first time. These are perhaps the largest close-coupling calculations carried out to date; yet we estimate that future relativistic calculations will be upto an order of magnitude more expensive in terms of computing resources and may not be feasible without massively parallel machines.

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