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Spectroscopic diagnostics of low-ionized iron-peak elements. 
Electron-impact excitation of Ni$^{3+}$ and photoionization of Ni$^{2+}$

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1 INTRODUCTION

The spectra of lowly ionized iron-peak elements such as (Fe$^{q+}$, Co$^{q+}$, Ni$^{q+}$, $q = 0$–$3$) are vitally important in astronomical observation. In particular, Ni$^{3+}$ being iso-electronic with Fe$^{+}$ should produce many of the same diagnostic lines, which have extensively been studied previously by Pradhan & Berrington (1993), Zhang & Pradhan (1995), Ramsbottom et al. (2005, 2007), and Ramsbottom (2009). Comparisons of these lines using the results of the present paper shall be made. The Fe-peak elements provide some of the most abundant species created inside the stars, and they emit at ultraviolet wavelengths, making them dominant contributors to the opacity of the interstellar media under certain conditions.

From an atomic physics perspective, the half-open d-shell nature of many of these systems inevitably leads to target descriptions involving between 20 and 30 configurations if spectroscopic accuracy is to be approached. The $N + 1$-electron collisional calculation, whether it be excitation or photoionization expands to target descriptions involving between 5000 and 7000 levels with the associated cost of calculating over $10^9$ Racah angular coefficients. Only with the development of the current suite of codes, including multiple layers of hyper-threaded parallelism, the Hamiltonian formation being the most critical part, have we been able to make progress on these type of systems. It has enabled us to provide comprehensive data sets that include every excitation and de-excitation for electron-impact excitation (EIE) or photoionization (PI), not just from the ground state, but as well from every excited state.

ABSTRACT

The spectra from Fe-peak elements may be used to determine the temperature and density of various astrophysical objects. Determination of these quantities is underpinned by the accuracy and the comprehensiveness of the underlying atomic structure and collisional calculations. In the following paper, we shall focus specifically on Ni$^{4+}$ lines associated with transitions amongst several low-lying levels. We shall employ modified versions of the parallel Dirac R-matrix codes, considering both electron-impact excitation of Ni$^{3+}$ and the photoionization of both the ground and excited states of Ni$^{2+}$. We produce high-quality data sets for both processes, and using these data, we calculate line ratios relevant for plasma diagnostics of temperature and density.

Key words: atomic data – opacity – techniques: spectroscopic.
large-scale (LS) resolved models, comprehensive level resolved calculations involving several hundred states remain to be calculated for the near-neutral Fe-peak elements.

Currently within the literature, some of the following works represent historical attempts at calculating lowly ionized iron peak elements, namely Pradhan & Berrington (1993), Zhang & Pradhan (1995) calculated the EIE of Mn-like Fe$^+$ in an LS coupling formalism. Their close-coupling (CC) expansion employed only three configurations, resulting in 38 LS terms, enabling transitions only among the ground and first excited configurations. We appreciate the limitations of this small model, and also that it has taken another 20 yr to include 20 more configurations in the configuration-interaction (CI) description of our present model. Ramsbottom et al. (2005, 2007); Ramsbottom (2009) also within an LS-coupling framework made a succession of calculations with a progressively better target description. They ultimately included orbitals up to $n = 4$ resulting in a total of 113 LS terms in their close coupling (CC) expansion. Other set of works for other isoelectronic sequences of low-ionized iron peak elements includes the one of Zhang & Pradhan (1997) for EIE of Fe$^{3+}$, and Bautista (2004) for EIE of Ni$^+$. Although not as dominant as iron, nickel lines are also used for diagnostic and modelling of astrophysical plasmas. Mazzali et al. (2001) performed several models for type Ia supernovae concluding that nickel abundance can affect its brightness and decline rate. Years later, nickel lines were observed in the remnant of the supernova 1987A by McCray & Fransson (2016). Furthermore, Werner, Rauch & Kruk (2018) subsequently used the absorption features in white dwarf atmospheres produced by iron-peaks elements to model the metal abundances. Opacity data are necessary for any kind of simulation work so the demand for comprehensive data sets is almost insatiable. One such example of these simulations is the work of Sánchez, Alfaro & Pérez (2007). They used models dependent upon opacities and the known optical depths of the interstellar clouds to determine its fractal dimension. Another example is the work of Moravveji (2016), whose opacity simulations, comparing measured to measured spectra, concluded that nickel ions produce an enhancement of the opacity.

Opacity is also an important aspect of the CLOUDY software package (Ferland et al. 2017). CLOUDY is extensively used for the simulation of the spectra collected from interstellar clouds. For completeness, the present work will investigate the Mn-like ion Ni$^{3+}$ for two important processes: the EIE and the PI of its parent ion Ni$^{2+}$. We employ a heavily modified parallel version of the fully relativistic Dirac atomic R-matrix code (DARC; Norrington & Grant 1987; Ballance & Griffin 2004). We include 23 configurations in the CI expansion. This expansion leads to a total of 6,841 relativistic levels. From that total, we reduce the CC expansion to include the first 262 levels, this reduction of the basis set may lead to pseudoerrors, and we have to take in account this fact when analysing the final collision strengths. Ni$^{3+}$ is a low-ionized intermediate-mass ion, therefore there is the expectation that relativistic effects will not be large, especially for valence-shell electrons. One might argue on theoretical grounds that a semirelativistic formalism, or even a non-relativistic one, would lead to acceptable results with considerably less computational effort. However, the current multilevel parallelism of the parallel DARC suite of codes, whilst being more computationally intensive is currently considerably more efficient than Breit-Pauli or ICFT semirelativistic versions.

Table 1. Configuration list included in the atomic structure calculations.

<table>
<thead>
<tr>
<th>Even parity</th>
<th>Odd parity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Core: $1s^2, 2s^2$</td>
<td>$1s^2$</td>
</tr>
<tr>
<td>$3p^6 3d^6 4s$</td>
<td>$3p^6 3d^6 4s$</td>
</tr>
<tr>
<td>$3p^6 3d^6 4s$</td>
<td>$3p^6 3d^6 4s$</td>
</tr>
<tr>
<td>$3p^6 3d^6 5s$</td>
<td>$3p^6 3d^6 5s$</td>
</tr>
<tr>
<td>$3p^6 3d^6 6s$</td>
<td>$3p^6 3d^6 6s$</td>
</tr>
</tbody>
</table>

Over the last few years considerable effort has been made by the group at Queen’s University Belfast in refactoring codes, specifically in terms of memory management. The last versions of the DARC code are viable, factoring in hardware limitations, to handle thousands of target states in the CI expansion.

The remainder of the paper is organized as follows: in Section 2 we give our description of the atomic structure; in Section 3 we describe the CC method used to obtain the EIE collision strengths and subsequent effective collision strengths as well as the PI cross-sections; in Section 4 we show and discuss the results; in Section 5 we perform a simple collision-radiative model to test the diagnostics predicted with present collision rates in relation to Fe II work; and in Section 6 we discuss the conclusions of the work. Atomic units are used unless otherwise specified.

2 STRUCTURE

We use the General-purpose Relativistic Atomic Structure Package (GRASP) (Dyall et al. 1989; Parpia, Fischer & Grant 1996) to determine the best possible atomic structure within a Dirac–Coulomb framework. The resulting radial orbitals from this Multiconfiguration Dirac-Fock (MCDF) method are defined on an exponential radial grid and they are employed subsequently in the EIE calculation.

In our CI expansion we permute the 25 electrons of the Mn-like Ni target within the configurations given below. Thirteen non-relativistic orbitals, namely the 1s, 2s, 2p, 3s, 3p, 3d, 4s, 4p, 4d, 5s, 5p, 6s, 6p are transformed into their relativistic counterparts within GRASP. To optimize the CI expansion and to accelerate the MCDF process, we follow several steps, validating our results against the recommended values of the NIST atomic spectra data table (Sugar & Corliss 1985; Kramida et al. 2018) where available. In our first step, we included the ground state configuration Ne 3s$^2$ 3p$^6$ 3d$^6$ 4s, and all possible one-electron excitations 3s$^2$ 3p$^6$ 3d$^6$ nl. This simple expansion led to a first approximation of the one-electron wave functions. Additional configurations only slightly refine the core orbitals up to the 3s, but do help the convergence of the valence orbitals. With each iteration we check the updated excitation energies of the first 50 levels with the recommended data of NIST, with the goal of a compact but accurate basis. The results of our final 23 configuration model are listed in Table 1, though we here only provide a representative sample of the possible 6,841 relativistic levels, the Supporting Information shall be more comprehensive. Comparing our calculated excitation energies with respect to the ground level with the recommended values of NIST we find our largest deviation in the order 12 per cent, and an average deviation of 3.3 per cent. This deviation is quite acceptable considering the complexity of the system and comparisons with the previous works for Mn-like
Fe of Ramsbottom et al. (2005), whose largest deviation was order 15 per cent in LS coupling, and the one of Pradhan & Berrington (1993), order 25 per cent.

For a further comparison and to quantify the uncertainty in the atomic structure we performed a second independent calculation using a different atomic structure code. The AUTOSTRUCTURE programme (Badnell 2011) code serves this purpose. AUTOSTRUCTURE provides non-relativistic radial wave functions from a Thomas–Fermi–Amaldi potential for the 1s to 6p orbitals. The subsequent Breit–Pauli Hamiltonian includes the relativistic terms as a first-order perturbations: mass-velocity, spin-orbit, and Darwin. We neglect the second-order perturbation terms spin–spin, orbit–orbit, and spin-other-orbit. To determine the $\lambda_{\text{ad}}$, or scaling parameters within the TFA model potential, we variationally determine them from minimization of the absolute Hamiltonian energy. For a balanced comparison with the GRASP and DARC calculations and in order to minimize the differences in atomic structure, we keep both problems as similar as possible. In that regard, we include in the CI expansion of the AUTOSTRUCTURE model exactly the same configuration set as that in GRASP. After performing the minimization process we obtained the values of $\lambda_{\text{ad}}$ shown in Table 2. In the Supporting Information, we show the energies obtained with AUTOSTRUCTURE for a complete comparison with the ones obtained with GRASP and in Table 3 we show here the lowest energy 50 levels. The level energies obtained with AUTOSTRUCTURE deviate slightly further from the recommended values of NIST than the ones obtained with GRASP. The maximum deviation is of the order of 15 per cent, again larger than the GRASP one.

To perform the CC integration including all the 6 841 levels obtained in the atomic structure is beyond the capabilities of existing workstations and even supercomputers. Consequently, we have selected the lowest excited 262 levels for the CC expansion. For analysis that favours the ground state and first few metastable states the completeness of this CC expansion is acceptable.

The Supporting Information will present a table of oscillator $f$ strengths and Einstein spontaneous emission coefficients $A$ for all the transitions between the 262 lowest excited levels. We show the values obtained with both methods GRASP and AUTOSTRUCTURE. This comparison gives an idea of the consistency for energies and transition probabilities for both atomic models. We also compare our results for the Einstein $A$-coefficients with previous theoretical calculations in the literature from Hansen, Raassen & Uylings (1984). Unfortunately, to the best our knowledge there are no experimental data available in the scientific literature for Ni$^{3+}$ to compare with.

Finally, to perform the scattering calculation we have shifted our calculated energies for the levels included in the CC expansion to the observed values of the NIST database. Doing so we make sure that the calculated wave lengths for the transitions will fit exactly with the observed ones, which is the requested for proper modelling of the astrophysical objects. In the NIST database, Ni$^{3+}$ has some missing energy levels for the highly excited states. Therefore, in those cases we have shifted our theoretical values by the difference with respect to the known NIST levels. We compare our final results using the shifted target energies with the unshifted ones as a test of accuracy.

3 SCATTERING AND PHOTOIONIZATION PROCESSES

We use an R-matrix formalism (Hummer et al. 1993; Berrington, Eissner & Norrington 1995). In the inner region, we use the fully relativistic DARC code (Ait-Tahar, Grant & Norrington 1996; Norrington & Grant 1981, 1987) to get the stationary solutions of the $N+1$ electron atom. We calculate the $N+1$ wave functions by diagonalization of the $N+1$ electron Hamiltonian. In addition, we calculate the dipole momentum matrices for the relevant photoionization transitions. In the outer region, we use the parallel version of the STGF programme to calculate the EIE collision strengths $\Omega$, and the radiative damped version PSTGF/DAMP for the photoionization cross-sections.

We calculate the photoionization cross-sections from several initial states of the parent ion Ni$^{3+}$. These levels are the relevant ones for an opacity model. With the available computational resources it is absolutely impossible to include in the CC expansion all the 6 841 levels calculated with the previous described CI expansion in GRASP. To have a reasonable accuracy in the calculation compatible with an affordable computation cost we have selected the 262 levels with the lowest energy for the CC expansion.

We use the same set of 262 levels to calculate the EIE of Ni$^{3+}$. We include partial waves with angular momentum up to $J = 36$.

3.1 Inner region

For our DARC calculation the R-matrix inner region radius is set to 59.52 au. We calculate the Hamiltonian matrices and the transition dipole momentum matrices. Including the first 262 levels of Ni$^{3+}$ target in the CC expansion we get a maximum of 1 818 channels in each $J^\pi$ symmetry.

For the photoionization calculation, we calculate partial waves with a total angular momentum of $J = 0$–5 and both parities. The lowest levels of the Ni$^{3+}$ ion have an angular momentum of $J = 0$–4 and even parity (see Kramida et al. 2018). Levels with higher angular momenta are very excited and they will rapidly decay to lower $J$ by an M1 or E2 transition. Levels with odd parity are very high in energy, the first one is the $3p^6 3d^3 4p^5 F_3^+$ with an energy of 1.0043 Ry relative to the ground state. They will be connected by an E1 transition to any lower level with even parity and their population will be zero in any astrophysical object. In addition, for each partial wave we calculate the dipole matrices with all their possible E1 couples.

For the EIE calculations, we need a more extended set of partial waves. We have calculated the energies and wave functions of the channels of all partial waves with an angular momentum of $J = 0$–36 and both parities plus a top-up.

3.2 Outer region

To calculate the photoionization cross-sections as a function of photon energy in Rydbergs, we utilize the parallel version of PSTGF/DAMP, a code which calculates the photoionization cross-sections utilizing the previously calculated bound-free matrix elements. The first serial version of STGF/DAMP was by Gorczyca & Badnell (unpublished material). The first stage is to determine the bound levels.

Table 2. Scaling parameters optimized by AUTOSTRUCTURE.

<table>
<thead>
<tr>
<th>$1s$</th>
<th>$1.42396$</th>
<th>$4s$</th>
<th>$1.04299$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2s$</td>
<td>$1.30959$</td>
<td>$4p$</td>
<td>$1.04410$</td>
</tr>
<tr>
<td>$2p$</td>
<td>$1.12342$</td>
<td>$4d$</td>
<td>$1.55730$</td>
</tr>
<tr>
<td>$3s$</td>
<td>$1.10133$</td>
<td>$5s$</td>
<td>$1.07620$</td>
</tr>
<tr>
<td>$3p$</td>
<td>$1.06211$</td>
<td>$5p$</td>
<td>$1.03593$</td>
</tr>
<tr>
<td>$3d$</td>
<td>$1.04845$</td>
<td>$6s$</td>
<td>$1.02930$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>$6p$</td>
<td>$1.01296$</td>
</tr>
</tbody>
</table>
of the \((N + 1)\)-electron system \(\text{Ni}^{2+}\) in the programme STGB (Seaton 1982; Berrington et al. 1987), which reads the wave functions for a specific partial wave in the inner region and determines its bound states. In our final calculated cross-sections, we shift the energies of the numerical \(\text{Ni}^{2+}\) levels to fit exactly the ionization potential with the values tabulated in NIST data basis. Hence, the threshold of the cross-sections fit exactly with the ionization potential of the initial state.

We split the energy range into two regions. In the low-energy region we adopt a fine energy mesh of \(1.5 \times 10^{-5}\) z\(^2\) Ry, being \(z = 3\) the charge of the final ion, to properly resolve the resonance structures converging on to the target thresholds. A linear grid with a total of 40 000 energy points was included up to the excitation energy of the last level included in the CC expansion. Above this threshold resonances are not present and the cross-sections are smoother, hence a coarser mesh of \(3 \times 10^{-5}\) z\(^2\) Ry was utilized. For higher
photon energies, above the excitation of the last included level in our CC expansion, 5.5 Ry in our case, there are more possible processes present in nature, for example the ionization with a final level which is not included in our CC expansion, or double ionization. Higher excited states, for example excitations $3s^{-1}$, while included in the CI expansion, are not included in the CC. Due to the limitation of our CC expansion, these processes cannot be reproduced by our model. Hence, present results are valid for a maximum photon energy of 5.5 Ry, approximately twice the ionization energy of Ni$^{2+}$ from its ground level.

For the EIE evaluation the parallel version of the STGF undamped package (Seaton et al., unpublished material) was utilized in the outer region. PSTGF calculates the outer region wave function using a Numerov method and including the coupling in the outer region as a perturbation. PSTGF joins the calculated wave function with one in the inner region in terms of the $R$-Matrix method (Burke 2011). In
the outer region problem high angular momenta do not contribute to the resonance structures, hence we restrict the fine-mesh calculation to the low partial waves with \( J = 0–20 \) and adopt a fine energy mesh of \( 1.5 \times 10^{-5} \) \( z^2 \) Ry, \( z = 3 \) being the ion charge, we incorporate a total of 40,000 points in the low-energy region. At higher energies, above the threshold energy of the last level included in the CC expansion, there is no more resonance structure, so we use a coarser mesh of \( 3 \times 10^{-3} \) \( z^2 \) Ry. The higher angular momenta \( J = 21 – 36 \) do not contribute to the resonance structure, even for low energies, hence the coarse mesh listed above is sufficient in the whole energy range. Finally, to include the remaining angular momenta up to \( J \) infinity we perform a top-up procedure. For dipole allowed transitions we use the Burgess sum rule Burgess (1974) and for the non-dipole allowed transitions with non-zero infinite energy Born limit a geometric series Badnell & Griffin (2001).

As the selected CC expansion in the target is considerably smaller than the initial CI expansion, we expect pseudoresonances to ap-

Figure 2. Electron-impact excitation effective collision strengths \( \Upsilon \) of Ni\(^{3+} \) for a Maxwellian electron distribution.
where \( T \) are temperatures where Ni\(^{3+}\) pseudoresonances will not affect the effective collision strengths at \( \frac{N_{\text{Ni}^{3+}}}{\Omega} \) necessary to calculate \( k \) in the outer region with PSTGF and the infinite energy point. For the have calculated, using DARC, the infinite-energy limit for the electric of a large number of continuum states in the Hamiltonian inclusion of a large number of continuum states in the Hamiltonian computed and thus increase the corresponding size of the matrices beyond \( \frac{\Omega_{ij}}{\Omega_{ij}} \) and \( \frac{\omega_{ij}}{\omega_{ij}} \). Data from Garstang (1958).

**4 RESULTS**

### 4.1 EIE of Ni\(^{3+}\)

The collision strength (\( \Omega_{ij} \)) between an initial state \( i \) and a final state \( j \) is directly related to the electron-impact collisional excitation cross-section \( \sigma_{ij} \) by

\[
\sigma_{ij} = \frac{\pi \alpha_{ij}^2}{\omega_k k^2},
\]

where \( \omega_0 = 2J_i + 1 \) is the statistical weight of level \( i \) and \( \alpha_k^2 \) is the incident electron energy in Rydberg. In the majority of astrophysical plasmas the electron velocity distribution is Maxwellian for a certain temperature \( T \). Hence, to aid plasma modelling we have performed a convolution of the collision strength \( \Omega_{ij} \) in terms of the Maxwellian distribution to obtain the associated effective collision strengths \( \Upsilon_{ij} \).

\[
\Upsilon_{ij} = \int_0^\infty \Omega_{ij}(E_j) \exp \left( -\frac{E_j}{kT} \right) d\left( \frac{E_j}{kT} \right),
\]

where \( T \) is the Maxwellian electron temperature in \( K \), \( E_j \) is the final energy of the incident electron, and \( k \) is Boltzmann’s constant. For high temperatures the Maxwellian has a long tail and it is necessary to calculate \( \Omega_{ij} \) up to a suitably high energy. This requires the inclusion of a large number of continuum states in the Hamiltonian and thus increase the corresponding size of the matrices beyond the computation capabilities available. To alleviate this problem we have calculated, using DARC, the infinite-energy limit for the electric dipole transitions and interpolated \( \Omega_{ij} \) in the scaled Burgess–Tully domain Burgess & Tully (1992) between the last energy calculated in the outer region with PSTGF and the infinite energy point. For the present calculation we can therefore guarantee accuracy of the \( \Upsilon_{ij} \) up to the order of 5.5 Ry, equivalent to \( 2 \times 10^5 \) K. The temperature of maximum-abundance for Ni\(^{3+}\) is approximately \( 5 \times 10^4 \) K (Mazzotta et al. 1998; Bryans et al. 2006), which indicates that the present evaluation is sufficient to model and resolve the emission features of the Ni\(\text{IV} \) lines in the range of temperatures where Ni\(^{3+}\) is abundant. We have used the programme ADASEXI (Griffin & Badnell, unpublished material) to perform the convolution of the \( \Omega \) and calculated the Maxwellian \( T \). We create a level-resolved specific ion adf04 file to store all the relevant collision-radiative parameters. This ADf04 file can be used as standard input to usual collision-radiative modelling software, for example the ADAS series 2 (Summers 1994).

In the present work, we have computed collision strengths \( \Omega_{ij} \) and effective collision strengths \( \Upsilon_{ij} \) for the EIE of the Ni\(^{3+}\) ion for transitions between the lowest 262 levels, a total of 34 191 forbidden and allowed lines. The highest energy considered was 5.5 Ry, adequate when compared to the ionization energy of 4.037 Ry (Kramida et al. 2018). Above this ionization energy, the collision strengths follow an asymptotic behaviour and can be interpolated with the infinite energy limit point in the Burgess–Tully domain Burgess & Tully (1992).

In Fig. 1, we present the collision strength \( \Omega_{ij} \) for the EIE of some selected transitions of the Ni\(^{3+}\) ion. For all transitions we observe the expected series of resonances in the low-energy region converging on to the target state thresholds included in the CC expansion, and a background above this that depends on the type of transition considered. The most useful transitions for astrophysical diagnosis are the M1 transitions between the levels of the ground term. A peculiarity of the present system is that the first levels with odd parity are highly excited, the first one listed as level 65. As a consequence of this the electric dipole E1 allowed transitions from the ground term are paradoxically very weak in comparison with the other M1 and E2 transitions within the lower excited levels, in fact it is in this transition where both versions of the calculation, with shifted and unshifted target energies, disagree the most (pannel g). The cause of this disagreement is that in both versions of the calculation, the wave functions of the atomic states have not been modified, but the energies have, since in one of them they have been shifted to the recommended values of NIST. As a consequence, the line strengths \( S \) have the same value in both calculations. At high energies, the collision strengths are determined by the infinite energy point, which in the case of E1 transitions depends only on the value of \( S \), see Burgess & Tully (1992). In Fig. 1, it is appreciated that for the E1 transition 1–65 the collision strengths obtained using both versions disagree at low impact energies, but they converge at high ones.

We present in Fig. 2 the corresponding Maxwellian averaged effective collision strengths \( \Upsilon_{ij} \) for the same transitions depicted in Fig. 1, for a range of electron temperatures \( T_e = 10^3 - 10^6 \) K. Clearly evident is the strong enhancement of the collision rates due to the proper delineation of the Rydberg resonance features in the collision strengths. For the Supporting Information, we provide tables of the calculated effective collision strengths for all the 34 191 transitions between all levels of Ni\(^{3+}\). For non-Maxwellian modelling or for any application that requires the direct collision strengths we direct the reader to our public ftp server. We also refer to the OPEN-ADAS data base for the general ADf04 file.

As a convergence test we have compared different ADf04 files, in the first one we have included in the partial wave expansion

Notes. WL, wavelength in air (Å); A, Einstein spontaneous emission coefficient \( s^{-1} \); \( A[B] \) denotes \( A \times 10^6 \). Data from Garstang (1958).
angular momenta up to $J = 30$ and no top-up, in a second one we have added the top-up to the $J = 30$ expansion, and finally our recommended data with the partial wave expansion extended up to $J = 36$ plus top-up. The largest differences remain between the versions with and without top-up, in that case the average difference between all the transitions values 0.5 per cent. In particular for the E1 allowed transitions, the maximum difference reaches the 100 per cent, while for the forbidden transitions this maximum difference is of the order 10 per cent. If we add the top-up to the $J = 30$ expansion the differences reduce significantly, the average difference is reduced to the 0.02 per cent, and the maximum difference for the E1 transitions to the 33 per cent, and only in six E1 transitions is above the 10 per cent, these six transitions are between very excited states, above 100, and they are irrelevant for the modelling. It is clear the calculation is properly converged in terms of the expansion in partial waves once the top-up is added, expansion up to $J = 30$ and $J = 36$ produce equal results.

Figure 3. Photoionization cross-sections versus the photon energy for Ni$^{3+}$ from ground lowest excited initial states. Colour figure is available online.
4.2 Photoionization of Ni\textsuperscript{2+}

We have calculated level resolved photoionization cross-sections of Ni\textsuperscript{2+} from its 20 lowest energy levels for each J\textsuperscript{\pi} symmetry with J = 0–4 and even parity, to all the 262 lowest excited levels of Ni\textsuperscript{3+}. These cross-sections can be considered of high-quality for photon energies up to 5.5 Ry and can be used for any opacity model.

In Fig. 4, a test of convergence for the calculation is presented. We compare two calculations performed with the same atomic structure of the target. In the first one (black line) we included in the configuration basis set of the (N + 1)-electron system all the configurations derived from the addition of one extra electron into all the available orbitals included in the expansion to those configurations listed in Table 1 and with an expansion of the continuum including N\textsubscript{c} = 20 functions. In the second one (red line) the configuration set was reduced somewhat extracting from Table 1 the 3p\textsuperscript{5} 3d\textsuperscript{7} 5s, 3p\textsuperscript{5} 3d\textsuperscript{7} 5p, 3p\textsuperscript{5} 3d\textsuperscript{7} 6s, 3p\textsuperscript{5} 3d\textsuperscript{7} 6p, 3p\textsuperscript{6} 3d\textsuperscript{4} 4d\textsuperscript{2}, 3p\textsuperscript{6} 3d\textsuperscript{4} 4d, 3p\textsuperscript{6} 3d\textsuperscript{9} basis configurations to build the (N + 1)-electron system expansion, and with N\textsubscript{c} = 13 functions for the expansion of the continuum. Evidently, there is a very small difference between both calculations with respect to the background, the position of the resonances and their heights. We can be confident therefore that the present calculation has converged with regard to the target description, the size of the continuum basis and the mesh size adopted in the low-energy region. For higher photon energies above 5.5 Ry a similar guarantee of the accuracy of the cross-sections cannot be made due to the effect of additional excited levels which are not included in our CC expansion.

As a test of accuracy we investigate in Fig. 5 the relative difference of the photoionization cross-sections produced when the target levels are shifted to their exact observed positions or left unshifted as the ab initio values. Relative differences of

\[ \delta = \frac{\sigma_{\text{sh}} - \sigma_{\text{un}}}{\sigma_{\text{sh}}} \]  

(3)

where \( \sigma \) represents the convoluted cross-section with a Gaussian enveloping for several widths. The largest deviation occurs for the lower photon energies. For those energies the difference in the positioning of the resonances is the dominant contribution to the global error. For photon energies above 3 Ry, equivalent to wavelengths shorter than 303.76 Å, the relative deviation for the convolution with width 10\textsuperscript{−2} Ry remains below the 10 per cent level in almost the entire domain. At a photon energy of E = 3.81 Ry the deviation reaches a maximum of 28 per cent, just for a single resonance. For a convolution width of 10\textsuperscript{−3} Ry and photon energies above 3 Ry the relative difference remains below the 20 per cent threshold. We estimate the accuracy of the present data to be approximately 20 per cent in the worst case for wavelengths in the ultraviolet, above the ionization limit of Ni\textsuperscript{2+}.

5 MODELLING OF DIAGNOSTICS

With the calculated effective collision strengths for the EIE of Ni\textsuperscript{3+}, we have performed a collision-radiative model. We use the programme COLRAD, which calculates the line intensities from the radiative transition probabilities and effective collision strengths stored in the ADF04 file. For low densities, the only mechanism of population is the collisional excitation from the ground or a
Table 5. Ni IV line ratios used for plasma diagnostics.

<table>
<thead>
<tr>
<th>Transition 1</th>
<th>Transition 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levels</td>
<td>Levels</td>
</tr>
<tr>
<td>20–21</td>
<td>4D9/2–6D7/2</td>
</tr>
<tr>
<td>1–5</td>
<td>4F9/2–4F5/2</td>
</tr>
<tr>
<td>1–26</td>
<td>4F9/2–4D5/2</td>
</tr>
<tr>
<td>1–5</td>
<td>4F9/2–4D5/2</td>
</tr>
</tbody>
</table>

Note. WL, wavelength in vacuum, in Å.

Figure 6. Line intensity ratio $I_{λ1}/I_{λ2}$ versus electron temperature and density for some selected pairs of lines of Ni IV. Colour online.

Figure 7. Line intensity ratio $I_{λ1}/I_{λ2}$ versus electron temperature for a constant density of $d = 10^4$ cm$^{-3}$ for lines (5 519.2 Å)/(9 524.8 Å) of Ni IV (full line) and (8 617.0 Å)/(12 566.8 Å) of Fe II (dashed line). Vertical lines indicate the peak abundance temperature of each ion. Colour figure is available online.

metastable state, following radiative-decay cascade. In Table 5, we have selected four line ratios to check their validity as diagnostics. These transitions were considered in a previous calculation by Pradhan & Zhang (1993) for the isoelectronic ion Fe$^+$, and hence provide a benchmark for the current analysis.

The line intensity ratios are plotted in Fig. 6 as a function of electron temperature and density. The ratio between the lines 1–5 and 20–25 (5 519.2 Å)/(9 524.8 Å) provides very powerful diagnostics for the electron temperature $T$. It is density independent and varies significantly in the range of the peak abundance temperature. The ratio 20–21/1–2 (127 345 Å)/(84 055 Å) similarly has a region where it is independent of density but the range is significantly greater than the temperature of maximum abundance for the Ni$^3+$ ion. The ratio between lines 1–26 and 1–25 (821.0 Å)/(827.1 Å) is a very useful density diagnostic particularly for low-density plasmas, below $10^9$ cm$^{-3}$, in the range of the temperature of peak abundance.

Additional line ratios can be analysed using the present effective collision strengths and with a more refined collision-radiative model. We provide good-quality data to perform plasma modelling using Ni IV emission lines.

For Fe II, the equivalent wavelengths to the ratio between 1–5 and 20–25 are the lines of 8 617.0 Å and 12 566.8 Å. The ratio of these lines can give a good diagnostic for the plasma temperature if it is in the range of the peak abundance for Fe$^+$ of $1.3 \times 10^4$ K, see (Smyth et al. 2019). Combining these two line ratios for Fe II and Ni IV, we are able to determine with accuracy the electron temperature of the plasma in a wider range. In Fig. 7, we show the variation of these line ratios for the electron density of the Orion nebula.

6 CONCLUSIONS

We present high-quality atomic data for EIE of Ni$^3+$ and photoionization from the ground and metastable levels of Ni$^2+$. These data are essential for the interpretation of Ni IV lines collected from...
ground and satellite observations, as well as opacity due to Ni\textsuperscript{+} in interstellar clouds. A fully relativistic DARC treatment is adopted with a configuration interaction expansion of the 25-electron target Ni\textsuperscript{+} incorporating lowest 262 levels in the close coupling expansion of the target. For each of the two processes, we have performed two calculations, one using the calculated energies and atomic wave functions obtained within GRASP, and a second one replacing the calculated energies with the recommended data tabulated in the NIST database. For both processes, the differences between the two calculations performed were negligible with the background cross-section as well as the height and positioning of the resonance structures almost identical in both. Accuracy checks were performed throughout the analysis and we are confident that the present data represent the best available to date for use by the astrophysics and plasma physics communities.

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**SUPPORTING INFORMATION**

Supplementary data are available at MNRAS online.

energies.dat
radiative.dat
upsilon.dat

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