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EXPANDED IRON UTA SPECTRA—PROBING THE THERMAL STABILITY LIMITS IN AGN CLOUDS

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ABSTRACT

The Fe unresolved transition arrays (UTAs) produce prominent features in the ~15–17 Å wavelength range in the spectra of active galactic nuclei (AGNs). Here, we present new calculations of the energies and oscillator strengths of inner-shell lines from Fe xiv, Fe xv, and Fe xvi. These are crucial ions since they are dominant at inflection points in the gas thermal stability curve, and UTA excitation followed by autoionization is an important ionization mechanism for these species. We incorporate these, and data reported in previous papers, into the plasma simulation code Cloudy. This updated physics is subsequently employed to reconsider the thermally stable phases in absorbing media in AGNs. We show how the absorption profile of the Fe xiv UTA depends on density, due to the changing populations of levels within the ground configuration.

Key words: atomic data – atomic processes – galaxies: active – line: formation – methods: numerical – X-rays: galaxies

Online-only material: color figures, machine-readable tables

1. INTRODUCTION

X-ray observations of active galactic nuclei (AGNs) have revealed the existence of a broad absorption feature in the 15–17 Å wavelength range. This feature arises from an unresolved transition array (UTA) due primarily to 2p → 3d inner-shell absorption lines in iron ions with an open M shell (Fe i–Fe xvi), and was first identified by Sako et al. (2001) in the XMM-Newton spectrum of the quasar IRAS 13349+2438. Since then, the UTA has been detected in numerous AGN (see, for example, Longinotti et al. 2010; Lestinsky et al. 2009, and references therein). It is believed to arise in the warm, relatively low ionization, absorbing gas surrounding the central supermassive black hole in the AGN.

Behar et al. (2001) first noted the diagnostic potential of the UTA, and showed that their central wavelengths, absorption profiles, and equivalent widths can provide information on the ionization structure, kinematics, and column densities of the warm absorbing material in the AGN. These authors also produced atomic data required for the modeling of the UTA, including energy levels (wavelengths) and oscillator strengths for the 16 iron charge states Fe i through Fe xvi, calculated with the multiconfiguration, relativistic HULLAC computer code (Bar-Shalom et al. 2001). Since then, several authors have calculated wavelengths and oscillator strengths for the UTA. These include Kisielius et al. (2003), who produced atomic data for Fe xv and Fe xvi using the configuration interaction code C1V3 (Hibbert 1975) with the inclusion of relativistic effects by adding Breit–Pauli operators to the Hamiltonian (Hibbert et al. 1991). More recently, Gu et al. (2006) calculated results for Fe vi through Fe xvi employing second-order many-body perturbation theory (MBPT; Lingren 1974). Furthermore, Beiersdorfer et al. (2012) used the relativistic multi-reference Møller–Plesset perturbation theory to calculate the energy levels of Fe xvi, including those of the autoionizing levels with a hole state in the L shell. They obtained very good agreement between the calculated L-shell transition wavelengths and those from recent laboratory measurements.

Here, we present extended calculations of UTA energies and radiative transition rates for Fe xiv, Fe xv, and Fe xvi. These are combined with other improvements in the plasma simulation code Cloudy (Ferland et al. 2013) to generate a range of models of absorbing clouds in AGN. Calculations were performed with r6433 of the Cloudy development branch, and will be part of the 2013 release of the code. We also perform a thermal stability analysis to identify the thermally stable phases where clouds can exist, and show that Fe xiv, Fe xv, and Fe xvi probe the high ionization end of the warm stable phase.

2. NEW ATOMIC DATA FOR Fe xiv, Fe xv, AND Fe xvi TRANSITIONS

Previously, a comparatively small set of inner-shell photoexcitation data for Fe xv and Fe xvi was reported by Kisielius et al. (2003). In that work energy levels were identified, and electric dipole transition wavelengths, oscillator strengths, and transition probabilities determined. Specifically, transitions involving photoexcitation from the inner 2p shell to the outer n = 3 shell were investigated using the relativistic Breit–Pauli approach, and an assessment of the accuracy of the derived atomic data was given. In the present work, the data set has been significantly extended. First of all, we add line data involving the transitions from the 2s shell since their wavelengths lie within the same UTA range. Furthermore, we determine excitation (both from 2s and 2p shells) to the n = 3, 4, and 5 shells. Finally, photoexcitation data for the same type of transitions in Fe xiv ions are derived.

In our current work, we use the configuration interaction (CI) approximation for atomic calculations. Particularly, the well-established computer code C1V3 of (Hibbert 1975) is exploited to
determine multiconfiguration eigenfunctions and eigenvalues of considered energy levels. After this initial step, we derive atomic data such as electric dipole (E1) radiative transition wavelengths $\lambda$, oscillator strengths $f$, and transition probabilities (rates) $A$ for the transitions from inner $n = 2$ shell of Fe xiv, Fe xv, and Fe xvi ions. The adopted CI method is suitable for production of very accurate data, where the correlation effects play a vital role in theoretical data accuracy.

Data are generated in the relativistic Breit–Pauli approach (Hibbert et al. 1991) for all lines arising from 2s and 2p shell electron transitions to the $n = 3, 4$, and 5 shells. Combining these two approaches, we can reliably determine $LS J$-energy levels and, consequently, the E1 transition data for fine-structure levels. When CI wave functions for the fine-structure levels of the ground and excited states are determined, the electric dipole transition absorption oscillator strength $f$ is given by

$$f = \frac{2}{3} g_{ij}^{-1} \Delta E_{ij} S^2,$$

(1)

where $g_{ij} = 2J_i + 1$ is the statistical weight of the initial (ground) level, $\Delta E_{ij}$ is the energy difference between the upper and lower level, and $S$ is transition line strength (matrix element). The corresponding electric dipole line emission transition probability $A$ (in $s^{-1}$) is given by

$$A = 2.142 \times 10^{10} g_{ij}^{-1} (\Delta E_{ij})^3 S^2,$$

(2)

where $g_{ij} = 2J_j + 1$ is the statistical weight of the excited-state level $j$.

A more detailed description of the approximations used in the present work may be found in Kisielius et al. (2003), where inner 2p shell photoexcitation was considered for Fe xiv and Fe xvi. The approach applied in the current paper for a much wider set of lines and Fe ions is very similar. Kisielius et al. (2003) have considered in detail the accuracy of determined results by applying mainly the internal indicators such as convergence of transition wavelengths with increasing the size of CI wave function expansion or agreement between length and velocity forms of oscillator strength values. They have estimated that 2p − 3/2 transition wavelength accuracy was 0.2% whereas oscillator strength accuracy was assessed as 10% for the lines having $f > 5 \times 10^{-5}$. The conclusions on the accuracy of the atomic data made by Kisielius et al. (2003) are also applicable to the present results.

2.1. Fe xvi Lines

The ground level of the Fe$^{15+}$ ion is 1s$^2$2s$^2$2p$^6$3s$^2$3p$^2$1S$^2$. Its CI wave function expansion was chosen to include no more than two electrons excited from the 2p and 3s shells in the case of the lines arising from the 2p shell, and no more than two electrons excited from 2s and 3s shells for the lines arising from the 2s shell. The significant difference between the calculation of $n = 3$ and 4 lines lies in the fact that we use correlation radial orbitals 4s, 4p, 4d, 4f in the former case to improve the calculated wavelength accuracy (for more details see Kisielius et al. 2003). In the latter case, these orbitals represent real states while the configuration sets remain the same.

2.1.1. Excitation from the 2p Shell

Following this approach, the ground state CI wave function expansion for the transition arrays 2p − 3/2 and 2p − 4/2 was the same and included 2p$^6$3l,2p$^5$4l ($l = 0, 2$), 2p$^5$3s3p, 2p$^3$3d, 2p$^3$3d'4l, 2p$^3$4l'4l' ($l = 0, 1, 2$; $l' = 0, 1, 2, 3$) even-parity configurations, coupled to the $^2S$, $^2P$, $^4P$, $^4D$ LS-terms. This CI expansion generates 94 LSJ-levels with total orbital quantum number $J = 1/2$. Due to selection rules for E1 transitions, the excited configuration with the vacancy in the 2p shell can only have levels with $J = 1/2$ and $J = 3/2$. We use a CI wave function expansion consisting of odd-parity configurations 2p$^6$3p, 2p$^5$4p, 2p$^5$3l'3d, 2p$^3$3d'4l' ($l = 0, 1, 2$), 2p$^3$3s3d, 2p$^3$4l'4l' ($l = 0, 1, 2, 3$), 2p$^3$4l'4l' ($l = 0, 1, 2, 3$) coupled to the $^2S$, $^2P$, $^2P$, $^2D$, $^2D$, $^4F$ terms for $J = 3/2$. Consequently, we consider 93 levels with $J = 1/2$ and 147 levels with $J = 3/2$ for the excited configurations having electrons in the outer $n = 3$ and 4 shells and the vacancy in the inner 2p shell.

Very similar configuration sets were used to obtain atomic data for the transitions from the 2p shell to the outer 5l shell. The ground state wave function expansion included 2p$^6$nl ($n = 3, 4, 5$; $l = 0, 2$), 2p$^3$3s3p, 2p$^3$3p3d, 2p$^3$3l'3l' ($l = 0, 1, 2$; $l' = 0, 1, 2, 3$), 2p$^3$3l'Sl' ($l = 0, 1, 2, 3$; $l' = 0, 1, 2, 3$) even-parity configurations coupled to the $^2S$, $^2P$, $^2P$, $^2D$, $^4D$, $^4L$-terms. This yields 95 LSJ-levels having $J = 1/2$.

The same (as in $n = 3, 4$ case) terms for the excited configuration levels with $J = 1/2$ and $J = 3/2$ were considered, while the CI expansion included configurations 2p$^6$np ($n = 3, 4, 5$), 2p$^3$3l' ($l = 0, 1, 2$), 2p$^3$3s3d, 2p$^3$3l'Sl' ($l = 0, 1, 2$; $l' = 0, 1, 2, 3$) 2p$^3$4l'2 ($l = 0, 1, 2, 3$), 2p$^3$5s5d, 2p$^3$5p5j. For $J = 1/2$, the CI expansion included 94 levels whereas for $J = 3/2$ the number was 148. Levels arising from the configurations with 5g electrons were not included in the CI expansion because such lines are very weak compared to other $n = 5$ transitions.

2.1.2. Excitation from the 2s Shell

A similar approach for the configuration sets to that adopted for the 2p shell was taken when calculating atomic data for the lines corresponding to the transitions from the inner 2s shell to the outer $n = 3, 4$, and 5 shells. For the $n = 3$ and 4 lines, the ground state CI expansion included 56 levels with $J = 1/2$ from the configurations 2p$^6$nl ($n = 3, 4$; $l = 0, 2$) 2p$^3$3s3p, 2p$^3$3p3d, 2p$^3$3l'3l' ($l = 0, 1, 2$), 2p$^3$3l'Sl' ($l = 0, 1, 2, 3$), 2p$^3$4l'2 ($l = 0, 1, 2, 3$), 2p$^3$4s4d, 2p$^3$4p4f coupled to the $^2S$, $^2P$, $^4P$, $^4D$ terms. The upper-state configuration set consisted of 2p$^3$3p, 2p$^3$4p, 2p$^3$4f, 2p$^3$3s3p, 2p$^3$3p3d, 2p$^3$4l'4l' ($n = 3, 4$; $l = 0, 1, 2$; $l' = 0, 1, 2, 3$) configurations coupled to the terms $^2S$, $^2P$, $^4P$, $^4D$, $^4D$, $^6F$ for $J = 1/2$, levels, and to the $^2P$, $^2D$, $^4S$, $^4P$, $^4D$, $^4F$, $^4P$, $^6D$, $^6F$, $^G$ terms for $J = 3/2$. This selection produced 38 and 58 levels for $J = 1/2$ and $3/2$, respectively.

For the lines corresponding to the transitions from the inner 2s shell to the outer $n = 5$ shell, the ground state wave function CI expansion included configurations 2p$^6$nl ($n = 3, 4, 5$; $l = 0, 2$), 2p$^3$3s3p, 2p$^3$3p3d, 2p$^3$3l'3l' ($l = 0, 1, 2$), 2s2p$^6$3s3d, 2s2p$^6$3l'Sl' ($l = 0, 1, 2$; $l' = 0, 1, 2, 3$), 2s2p$^6$4l'2 ($l = 0, 1, 2$), 2s2p$^6$5s5d, 2s2p$^6$5p5j coupled to the terms $^2S$, $^2P$, $^4P$, $^4D$, $^4D$, $^6F$, producing 39 levels with $J = 1/2$. For $J = 3/2$, we considered the same terms as in the $n = 4$ lines case, producing 59 levels.

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2.1.3. Transition Data

We employ multiconfiguration wave functions determined after the Hamiltonian matrix diagonalization to calculate transition wavelengths $\lambda$, oscillator strengths $f$, and transition rates $A$ for excitation from the inner $n = 2$ shell to the outer $n = 3, 4$, and 5 shells. Calculated line data cover the wavelength region $\lambda = 12.606 - 14.096$ Å for $2s - 3l$ transitions (a total of 17 lines), $\lambda = 10.145 - 11.221$ Å for $2s - 4l$ transitions (46 lines), and $\lambda = 9.545 - 10.333$ Å for $2s - 5l$ transitions (46 lines). Considering transitions from the $2p$ shell, the wavelength range was $\lambda = 13.594 - 17.246$ Å for the excitation of the $n = 3$ shell (45 lines), $\lambda = 11.316 - 12.934$ Å for the $n = 4$ shell (111 lines), and $\lambda = 10.525 - 11.689$ Å for the $n = 5$ shell (111 lines).

In Table 1, we present data for Fe xvi transitions from the $2s$ and $2p$ shells to $n = 3, 4$, and 5. We include only transitions with $f \geq 0.1$, whereas the more complete online version of the table contains lines having $f \geq 0.0001$. Weaker lines are not presented since they do not affect the ionization balance nor the final spectra. More details on data trimming are given in Section 3.2.

In Figure 1, we compare the strongest lines from our calculation with data derived using an MBPT (Gu et al. 2006), and with the Behar et al. (2001) results obtained using the multiconfiguration relativistic HULLAC computer package. The Behar et al. (2001) data are given for an $f$-value averaged wavelength $\lambda_{av}$ and the total oscillator strength $\sum f_{ij}$ of the line group, and therefore there is a sizable difference in the number of lines and in the values of $f$. Note that the point at $\lambda = 17.296$ Å represents transitions from the $2p$ to $3s$ shells which are not considered in our calculations. When comparing our results with those of Gu et al. (2006), one can see quite good agreement both for wavelengths and oscillator strengths. There is some systematic shift toward shorter wavelengths in our data, as discussed in Kisielius et al. (2003). In addition, a significant difference with Gu et al. (2006) is the appearance in our results of a group of lines at $\lambda = 11.5 - 12.5$ Å which represent transitions to $n = 4$ and $n = 5$ states. Some of these have large $f$-values and are important to include in atomic models.

The primary goal of the calculations presented here is to create an extensive set of inner-shell transition data to ensure that the total excitation and ionization rate is computed as accurately as possible. In a later section, we show how this new data change the ionization distribution of gas near an AGN. The line wavelengths we quote, which come from our atomic structure calculation, are no better or worse than the wavelengths quoted in previous studies. Indeed, the scatter in the wavelengths represents the uncertainty. Extensive sets of laboratory measurements of energy levels would be needed to improve the level energies and resulting wavelengths. Although a few experiments have been done (Brown et al. 2001; Simon et al. 2010), the extensive laboratory data needed to significantly improve the line wavelengths do not now exist.

2.2. Fe xvi Lines

The ground level of the Fe$^{15+}$ ion is $1s^22s^22p^63s^2$. Its CI wave function expansion was chosen to include no more than two electrons virtually excited from the $2p$ and $3s$ shells in the case of the lines arising from the $2p$ shell, and no more than two electrons excited from the $2s$ and $3s$ shells for the lines arising from the $2s$ shell. We adopt the same configuration set both for the $2p - 3l$ and the $2p - 4l$ line calculations, but different radial orbital sets are used. For the $n = 3$ line calculation, we employ the correlation radial orbitals $4s, 4p, 4d,$ and $4f$, whereas the corresponding radial orbitals represent real states in the $n = 4$ line calculations. This method contributes to an increased accuracy of the calculated wavelengths for $n = 3$ lines which are much stronger than the $n = 4$ ones.

### Table 1

<table>
<thead>
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<th>$n_i l_i$</th>
<th>$n_j$</th>
<th>$g_i$</th>
<th>$g_j$</th>
<th>$\lambda(\text{Å})$</th>
<th>$f_{ij}$</th>
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<tr>
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<td>4</td>
<td>14.996</td>
<td>0.697</td>
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<tr>
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<td>2</td>
<td>4</td>
<td>12.336</td>
<td>0.230</td>
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<tr>
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<td>4</td>
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<td>13.971</td>
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**Notes.** Column $n_i l_i$ denotes the initial shell, $n_j$ denotes the final shell for optical electron transition, $g_i$ and $g_j$ denote the statistical weights for the initial and final levels, $\lambda$ denotes the line wavelength (in Å), and $f_{ij}$ denote the absorption oscillator strength.

(This table is available in its entirety in a machine-readable form in the online journal. A portion is shown here for guidance regarding its form and content.)

![Figure 1](image-url)
2.2.1. Excitation from the 2p Shell

For excitation from 2p to the n = 3 and 4 shells, a CI wave function expansion was made of the 2p^63l^2 (l = 0, 1, 2), 2p^63s3d, 2p^63s^23p, 2p^63p^3, 2p^63p^2d, and 2p^63s^23p^2d configurations containing two or three electrons within the n = 3 shell. The even-parity configurations 2p^63l^4l′ (l = 0, 1, 2; l′ = 0, 1, 2, 3), 2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) of this ion. In the same way as for the lower shell, the odd-parity configurations 2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) for the upper (odd) state of this ion. The CI wave function expansion consists of the 2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) wave function expansion for the upper (odd) state of this ion. In the same way as for the lower shell, we consider the 1S^o, 3P^o, and 3D^o terms which produce the J = 0 fine-structure levels. Construction of such a basis produces 408 configuration state functions (CSFs) in the CI wave function expansion for the lower state of the Fe^{14+} ion.

Due to the electric dipole transition selection rules, we have to consider fine-structure levels with J = 1 in the upper (odd) state of this ion. In the same way as for the lower (even) state, we construct three subsets of configurations in wave function expansion for the upper (odd) state. The CI wave function expansion consists of the 2p^63s3p, 2p^63p3d, 2p^63s^23d, 2p^63s^23p^2, 2p^63p^2d, and 2p^63d^3 odd-parity configurations which have two or three electrons in the n = 3 shells. The odd-parity configurations 2p^63l^4l′ (l = 0, 1, 2; l′ = 0, 1, 2, 3), 2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) with two electrons excited to the n = 4 shell. In the case of inner 2p shell excitation, we consider the 1S^o, 3P^o, and 3D^o terms which produce the J = 0 fine-structure levels. Construction of such a basis produces 408 configuration state functions (CSFs) in the CI wave function expansion for the lower state of the Fe^{14+} ion.

In this construction of the CI wave function basis, we restricted our set by allowing no more than two electrons in the excited n = 4 shell. Here, we consider the 3S^o, 1P^o, 3P^o, 5P^o, 3D^o, 5D^o, and 3F^o terms which can produce fine-structure levels with J = 1. As a result, we have the CI wave function expansion with 1094 CSFs for the upper states of the Fe^{14+} ion.

A very similar configuration set was applied when considering 2p – 5l transitions. In this instance, the configurations with n = 4 electrons were replaced by configurations with n = 5. The only significant difference was the addition of the 2p^63l^4l′ (l = 0, 1, 2; l′ = 0, 1, 2, 3, 4, 5) for CI expansion of the lower state of Fe^{14+}, and the 2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) odd-parity configurations for the excited state. Configurations with 5g electrons were not included in the CI wave function expansion because transitions to such configurations are very weak. The total number of fine-structure levels of J = 0 in this case was 413 for the lower configuration and 1107 for the upper configuration levels with J = 1.

2.2.2. Excitation from the 2s Shell

While determining the transition array 2s – 3l, 4l lines, for the CI expansion we have adopted the same subset (as in 2p – 3l, 4l case) of configurations with closed 2s^2 and 2p^6 electron shells and complemented it with configurations having one electron virtually excited from the 2s shell. Specifically for the ground level, the even-parity configurations 2s2p^63s3d, 2s2p^63p3d^2, 2s2p^63p^23p^2, 2s2p^63s^23p^2, 2s2p^63s^23d, 2s2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) and 2s2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) were included in our CI wave function expansion. These configurations were coupled in the 1S^o, 3P^o, and 3D^o non-relativistic LS-terms and produced 165 fine-structure levels with total orbital quantum number J = 0.

The construction of the configuration set for the upper state is based on the same principles. We employ configurations with closed 2s^2 and 2p^6 electron shells from the 2p – 3l, 4l array calculation and add the configurations with an open 2s shell. Following this method, we include the configurations 2s2p^63s^23p, 2s2p^63p3d^2, 2s2p^63p^2, and 2s2p^63s^23d, which can produce the 2s electron virtually excited to the n = 3 shell, the odd-parity configurations 2s2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) and 2s2p^63l^4l′ (l = 0, 1, 2, 3, 4, 5) with two electrons excited to the n = 4 shell. In the case of inner 2s shell excitation, we consider the 1S^o, 3P^o, and 3D^o terms which produce the J = 0 fine-structure levels. Construction of such a basis produces 408 configuration state functions (CSFs) in the CI wave function expansion for the lower state of the Fe^{14+} ion.

2.2.3. Transition Data

Similarly to the Fe xvi case, we obtained multiconfiguration wave functions to produce photoexcitation data for the Fe xiv UTA lines. The calculated line data cover the wavelength region x = 11.592–14.199 Å for 2s – 3l transitions (a total of 35 lines), x = 9.620–11.423 Å for 2s – 4l transitions (147 lines), and x = 9.545–10.333 Å for 2s – 5l transitions (147 lines). Considering transitions from the 2p shell, the wavelength range was x = 12.475–16.266 Å for the excitation of the n = 3 shell (88 lines), x = 10.630–13.244 Å for the n = 4 shell (384 lines), and x = 9.998–12.017 Å for the n = 5 shell (384 lines).

In Table 2, we present a sample of the line data for Fe xv. Transition wavelengths x and oscillator strengths f_ji for excitation from the 2s and 2p shells to the outer n = 3, 4, and 5 shells are presented. Only a small portion of lines with f ≥ 0.04 is included, with the more complete online version of the table containing all lines which have f ≥ 0.0001. It is evident from Table 2 that the lines representing transitions to the n = 3 and 4 shells are sufficiently strong as to affect spectral formation, and therefore they cannot be excluded from UTA line data sets.

Figure 2 compares the strongest lines from our calculations with the results derived using the MBPT approach (Gu et al. 2006) and with the relativistic data from Behar et al. (2001). In general, the agreement between all three sets of data is satisfactory. There is a systematic shift toward shorter wavelengths in our data set as noted for the Fe xvi lines.

2.3. Fe xiv Lines

The ground configuration of the Fe^{14+} ion is 1s^22s^22p^63s^23p. It has two fine-structure levels, ^1P^o_{1/2} and ^3P^o_{3/2}, separated by
et al. 2003), for photoexcitation from the 2p and 2s shells of the ground configuration to the levels of the excited \( n = 3 \) configurations. Furthermore, for the excitation of the \( n = 4 \) and \( n = 5 \) lines, we adopt only real radial orbitals to describe 4l and 5l (\( l = 0, 1, 2, 3 \)) electrons.

2.3.1. Excitation from the 2p Shell

We employ the same configuration set both for calculations of 2p - 3l and 2p - 4l transition data. The CI expansion for the lower state includes configurations which have one or two electrons virtually excited from the \( n = 3 \) shell. These are complemented with configurations which have the 2p electron moved to the \( n = 3 \) shell. The CI wave function expansion for the ground state was composed of the configurations with three or four electrons in \( n = 3 \) shell \( 2p^63s^23p, 2p^63p3d^2, 2p^63p^23d, 2p^63s^23p^2, 2p^63p^23d^2, 2p^63s^33d, 2p^63p^4, 2p^53d^2, \) and \( 2p^53s^23p^2 \), the odd-parity configurations with one electron in virtually excited \( n = 4 \) shell \( 2p^63l^24l' \) (\( l = 0, 1, 2; l' = 1, 3 \)) and \( 2p^63l^23l'4l'' \) (\( l = 0, 1; l' = 1, 2; l'' = 0, 1, 2, 3 \)), and the odd-parity configurations with two electrons in \( n = 4 \) shell \( 2p^63l^24l'' \) (\( l = 0, 1; l' = 1, 2; l'' = 0, 1, 2, 3 \)).

In the case of the \( J = 1/2 \) levels, we consider the \( 2S, 2P, 4P, 4D, 6D, \) and \( 6F \) odd-parity LS-terms making up 336 CSFs in the CI wave function expansion, while for the \( J = 3/2 \) fine-structure levels, the odd-parity LS-terms \( 2P, 2D, 4S, 4P, 4D, 6P, 6F, \) and \( 6G \) produce 553 CSFs.

For the upper state levels, the CI wave function expansion for the 2p - 4l lines calculation is constructed from even-parity configuration sets. The configurations with electrons in \( n = 3 \) shell \( 2p^63s^23d, 2p^63s^23p^2, 2p^63s^23d^2, 2p^63p^23d^2, \) and \( 2p^63s^23p^2 \) the even-parity configurations with one electron in \( n = 4 \) shell \( 2p^63l^24l' \) (\( l = 0, 1, 2; l' = 0, 2 \)) and \( 2p^63l^23l'4l'' \) (\( l = 0, 1; l' = 1, 2; l'' = 0, 1, 2, 3 \)) and the even-parity configurations with two electrons virtually excited into the \( n = 4 \) shell, namely, \( 2p^63l^24l'' \) (\( l = 0, 1; l' = 0, 1, 2; l'' = 0, 2 \)) and the even-parity configurations with a vacancy in the inner 2p shell: \( 2p^53s^23d, 2p^53s^23d^2, 2p^53s^23p, 2p^53s^23d^2, 2p^53s^23d^2, 2p^53s^23d^2, 2p^53s^23d^2, 2p^53s^23d^2 \) (\( l = 0, 1; l' = 1, 2; l'' = 0, 1, 2, 3 \)) and \( 2p^53l^24l'' \) (\( l = 0, 1; l' = 0, 1, 2, 3 \)), \( 2p^53l^24l'' \) (\( l = 0, 1, 2; l'' = 0, 1, 2, 3 \)), \( 2p^53l^24l'' \) (\( l = 0, 1; l' = 1, 2; l'' = 0, 1, 2, 3 \)), \( 2p^53l^24l'' \) (\( l = 0, 1, 2; l'' = 0, 1, 2, 3 \)), \( 2p^53l^24l'' \) (\( l = 0, 1; l' = 1, 2; l'' = 0, 1, 2, 3 \)).

These configurations are bounded to the \( 2S, 2P, 4P, 4D, 6P, 6D, \) and \( 6F \) LS-terms for \( J = 1/2 \) and produce 4099 CSFs in the CI wave function expansion. The terms \( 2P, 2D, 4S, 4P, 4D, 6P, 6F, 6G, 6H \) and \( 6P \) generate a total of 7049 fine-structure levels for \( J = 3/2 \). For the total angular momentum \( J = 5/2 \), the above configurations are bound to the \( 2D, 2F, 2P, 2D, 4F, 6G, 6S, 6P, 6D, 6F, 6G, \) and \( 6H \) non-relativistic terms and produce 8211 levels.

The CI expansion for the lower state in the case of 2p - 5l consists of configuration sets similar to those used in 2p - 3l, 4l lines calculation. However, the four configuration complexes with 4l (\( l = 0, 1, 2, 3 \)) electrons are replaced by the configurations with \( 5l \) electrons. Furthermore, the CI wave function expansion is extended by an additional set of the odd-parity configurations \( 2p^63l^24l' \) (\( l = 0, 1, 2; l' = 1, 3 \)) and \( 2p^63l^23l'4l'' \) (\( l = 0, 1; l' = 1, 2; l'' = 0, 1, 2, 3 \)) with one electron in an outer \( n = 4 \) shell. The same (as in case of transitions to \( n = 3, 4 \)) non-relativistic LS-terms produce 359
CSFs in the CI wave function expansion for $J = 1/2$, and 591 CSFs for $J = 3/2$.

Similarly, the configuration sets with 4l electrons are replaced by the configuration sets with 5l electrons for the upper states. This set is extended with the even-parity configurations having one electron in the $n = 4$ shell: $2p^63l^24l'$ ($l = 0, 1, 2; l' = 0, 2$) and $2p^63l'3'l''4l''$ ($l = 0, 1; l'' = 1, 2; l'' = 0, 1, 2, 3$). As in the previous case, even-parity LS-terms produce 4326, 7421, and 8612 CSFs in the CI wave function expansion for total angular momentum $J = 1/2, 3/2,$ and 5/2, respectively.

### 2.3.2. Excitation from the 2s Shell

For the lines representing transitions from the inner 2s shell to the valence $n = 3$ or $n = 4$ shells, the CI wave function expansion configurations for the lower state consists of the same configuration complexes for both cases. The first are configurations with outer electrons in $n = 3$ shell $2p^63s^33p$, $2p^63s^33p^2$, and $2p^63s^23p^3$ and those with 2s vacancy $2s2p^63s^23s^3d$, $2s^2p^63s^23p^3d^2$, $2s^2p^63s^23p^3d^3$, and $2s^2p^63s^23p^3d^4$. These configurations are complemented with odd-parity configurations with one and two electrons in the 4l shell: $2p^63l^24l'$ ($l = 0, 1, 2; l' = 1, 3$), $2p^63l'3'l''4l''$ ($l = 0, 1; l'' = 1, 2, l'' = 0, 1, 2, 3$), $2p^63l'3'l''4l''$ ($l = 0, 1; l'' = 1, 2, l'' = 0, 1, 2, 3$), $2p^63l'3'l''4l''$ ($l = 0, 1, 2; l'' = 0, 1, 2, 3$), and $2p^63l's3d3d4l'$ ($l = 0, 2$).

In the case of the $J = 1/2$ levels, the above configurations are bound to the $2s^2, 2p^63s^33p$, $2p^63s^23p^2$, and $2p^63s^23p^3$ odd-parity LS-terms which yield a total of 660 CSFs in the CI wave function expansion. For the $J = 3/2$ levels represented by the $2p^6$, $2d$ $4s, 4p, 2D, 4F, 6p, 6D, 6F$, and $6G$ odd-parity LS-terms, the wave function expansion consists of 1102 CSFs.

The wave function expansion for the lower state in the case of the 2s – 5l transition array is constructed in a similar way by replacing all the configurations containing 4l ($l = 0, 1, 2, 3$) electrons with configurations containing 5l electrons. Furthermore, a complex of configurations with one electron in the 4l shell is added. This extends CI expansion by including odd-parity configurations $2p^63l^24l'$ ($l = 0, 1, 2; l' = 1, 3$) and $2p^63l'3'l''4l''$ ($l = 0, 1; l'' = 1, 2, l'' = 0, 1, 2, 3$). The same (as for the $n = 4$ lines) LS-terms are considered, and the CI wave function expansion gives rise to 716 and 1193 CSFs for the $J = 1/2$ and $J = 3/2$ levels, respectively.

While considering the upper state of Fe$^{13+}$, the CI wave function expansion for 2s – 5l, 4l lines is constructed of several configuration complexes, representing configurations with one or two electrons virtually excited from the ground configuration. We have included configurations $2p^63s^23d^2, 2p^63s^23d^02p^63d^03d^2, 2p^63d^2$, and $2p^63d^3$ with three outer electrons in $n = 3$ shell, the even-parity configurations $2p^63l^24l'$ ($l = 0, 1, 2; l' = 0, 2$), $2p^63l'3'l''4l''$ ($l = 0, 1; l'' = 1, 2, l'' = 0, 1, 2, 3$), and $2p^63l'3'l''4l''$ ($l = 0, 1; l'' = 0, 1, 2, 3$) with one electron in outer $n = 4$ shell, $2p^63l's3d4l'$ ($l = 0, 1, 2; l = 0, 1, 2; l'' = 1, 2, 3$) with two electrons in $n = 4$ shell. On top of that, we have extended the CI expansion by adding the even-parity configurations with a vacancy in the 2s shell: $2s^2p^63l^2l'^2$ ($l = 0, 1, 2; l' = 1, 2$), $2s^2p^63s^23p^23d^2, 2s^2p^63s^23p^3d^3, 2s^2p^63s^23p^4, 2s^2p^63d^4, 2s^2p^63l^23l''4l''$ ($l = 0, 1, 2, l'' = 1, 2, l'' = 0, 1, 2, 3$), $2s^2p^63l's3l''4l''$ ($l = 0, 1, 2, l'' = 0, 1, 2, 3$), $2s^2p^63l's3l''4l''$ ($l = 0, 1, 2, l'' = 0, 1, 2, 3$) $2s^2p^63l's3l''4l''$ ($l = 0, 1, 2, l'' = 0, 1, 2, 3$).

### Table 3

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Notes. Column $n_{li}$ denotes the initial shell, $n_j$ denotes the final shell for optical electron transition, $g_l$ and $g_j$ denote the statistical weights for the initial and final levels, $\lambda$ denotes the line wavelength (in Å), and $f_j$ denotes the absorption oscillator strength.

(This table is available in its entirety in a machine-readable form in the online journal. A portion is shown here for guidance regarding its form and content.)

### 2.3.3. Transition Data

Using the above methods to generate the CI wave function expansions, we obtained wave functions to produce photoexcitation line data for the Fe xiv ion. Calculated line data cover the wavelength region $\lambda = 10.571$–14.432 Å for 2s – 3l transitions (a total of 617 lines), $\lambda = 9.298$–11.707 Å for 2s – 4l transitions (3194 lines), and $\lambda = 8.890$–10.861 Å for 2s – 5l transitions (147 lines). For transitions from the 2p shell, the wavelength range was $\lambda = 12.341$–16.626 Å for excitation from the $n = 3$ shell (1343 lines), $\lambda = 10.141$–13.642 Å for the $n = 4$ shell (7425 lines), and $\lambda = 9.685$–12.438 Å for the $n = 5$ shell (8640 lines).

Table 3 lists the atomic data for the Fe xiv UTA lines. Transition wavelengths $\lambda$ and absorption oscillator strengths $f_j$ for excitation from the 2s and 2p shells to the outer $n = 3, 4,$ and 5 shells are given for lines with $f \geq 0.1$. A more extensive data set is presented in the online version of this.
Figure 3. Comparison of the wavelengths $\lambda$ and absorption oscillator strengths $f_{ij}$ for the strongest lines in Fe $^{14+}$. Dots represent our data, crosses are the results of Gu et al. (2006), and stars are those from Behar et al. (2001). The data at $\lambda = 15.5$ Å represent the $2p - 3d$ UTA, those at $\lambda = 14.3$ Å represent the $2s - 3p$ UTA, and those at $\lambda = 12.9$ Å represent the $2p - 4d$ UTA. The Behar et al. (2001) data are given for an $f$-value averaged wavelength $\lambda_{av}$ and the total oscillator strength $\sum f_{ij}$ of the line group. (A color version of this figure is available in the online journal.)

Table, where we provide results for all lines with $f > 0.0001$ (see Section 3.2 for more details about data trimming).

In Figure 3, we compare the strongest lines from our calculation with the results derived using the MBPT approach of Gu et al. (2006) and with the HULLAC relativistic data from Behar et al. (2001). There is reasonable wavelength agreement for all three sets, as found for Fe $^{16+}$ and Fe $^{15+}$. The only exception is the line with $\lambda = 16.033$ Å from Behar et al. (2001), with the larger values of oscillator strengths $\sum f_{ij}$ found by these authors explained in Section 2.1.3. A wavelength disagreement between our results and the MBPT data was also discussed in that section.

3. INCORPORATION INTO CLOUDY

3.1. Data Sources

The UTA data sets now used in Cloudy are summarized in Table 4. The Opacity Project (OP; Badnell et al. 2005) produced complete data, for the K shell as well as the L1 and L2 shells where appropriate, for ions with 12 or fewer bound electrons. These include the autoionization branching ratio which we consider in calculating the ionization rate. The OP data are marked “B” in this table. Gu et al. (2006) provide all stages of ionization for Fe but includes only excitations from the $2p$ shell, denoted by “G” in the table. These data sets have been included in Cloudy since shortly after their original publication dates.

The data presented in this paper are now being incorporated into Cloudy and are denoted by “K” in Table 4. The Opacity Project (OP; Badnell et al. 2005) produced complete data, for the K shell as well as the L1 and L2 shells where appropriate, for ions with 12 or fewer bound electrons. These include the autoionization branching ratio which we consider in calculating the ionization rate. The OP data are marked “B” in this table. Gu et al. (2006) provide all stages of ionization for Fe but includes only excitations from the $2p$ shell, denoted by “G” in the table. These data sets have been included in Cloudy since shortly after their original publication dates.

The newly computed data sets are quite extensive and include a large number of exceptionally weak lines. Figure 4 shows the absorption line oscillator strength $f_{l,u}$ plotted against the normalized sum of all lines with $f > f_{l,u}$. Careful examination of such data shows that the summed oscillator strength, which affects the UTA ionization rate, has converged to within 0.57%, 0.15%, and 0.08% of the total for Fe $^{13+}$, Fe $^{14+}$, and Fe $^{15+}$, when all lines with $f > 10^{-4}$ are included. Weaker lines have little effect upon the ionization rate because the line-center opacity is smaller than the continuous photoionization opacity. Accordingly, we only consider lines with $f \geq 10^{-4}$, although we retain all data for future flexibility. For the case of Fe $^{13+}$, this reduces the number of lines from 80894 to 754, for Fe $^{14+}$ from 2738 to 103, and for Fe $^{15+}$ from 595 to 137.

3.2. Data Trimming

The newly computed data sets are quite extensive and include a large number of exceptionally weak lines. Figure 4 shows the absorption line oscillator strength $f_{l,u}$ plotted against the normalized sum of all lines with $f > f_{l,u}$. Careful examination of such data shows that the summed oscillator strength, which affects the UTA ionization rate, has converged to within 0.57%, 0.15%, and 0.08% of the total for Fe $^{13+}$, Fe $^{14+}$, and Fe $^{15+}$, when all lines with $f \geq 10^{-4}$ are included. Weaker lines have little effect upon the ionization rate because the line-center opacity is smaller than the continuous photoionization opacity. Accordingly, we only consider lines with $f \geq 10^{-4}$, although we retain all data for future flexibility. For the case of Fe $^{13+}$, this reduces the number of lines from 80894 to 754, for Fe $^{14+}$ from 2738 to 103, and for Fe $^{15+}$ from 595 to 137.

3.3. Damping Parameters and Line Broadening

The Voigt function, which describes the line profile including both thermal and natural broadening, is given by

$$H(a, x) = \frac{a}{\pi} \int_{-\infty}^{\infty} \frac{\exp(-y^2)}{(x - y)^2 + a^2} \, dy, \quad (3)$$
Table 4

Data Sources used in the Default Setup of the Cloudy Code

| Ion | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 | 23 | 24 | 25 | 26 | 27 | 28 | 29 | 30 |
|-----|---|---|---|---|---|---|---|---|---|---|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| H   |   |   |   |   |   |   |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| He  |   |   |   |   |   |   |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Li  | B |   |   |   |   |   |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Be  | B | B |   |   |   |   |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| B   | B | B | B |   |   |   |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| C   | B | B | B | B |   |   |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| N   | B | B | B | B | B |   |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| O   | B | B | B | B | B | B |   |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| F   | B | B | B | B | B | B | B |   |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Ne  | B | B | B | B | B | B | B | B |   |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Na  | B | B | B | B | B | B | B | B | B |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Mg  | B | B | B | B | B | B | B | B | B | B |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Al  | B | B | B | B | B | B | B | B | B | B | B |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Si  | B | B | B | B | B | B | B | B | B | B | B | B |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| P   | B | B | B | B | B | B | B | B | B | B | B | B | B |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Cl  | B | B | B | B | B | B | B | B | B | B | B | B | B | B |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |    |
| Ar  | B | B | B | B | B | B | B | B | B | B | B | B | B | B | B |   |    |    |    |    |    |    |    |    |    |    |    |    |    |    |


where $x$ is the displacement from line center, measured in terms of the Doppler width $\Delta \nu_{Dop}$,

$$x \equiv \frac{\nu - \nu_0}{\Delta \nu_{Dop}}.$$  

(4)

The damping parameter $a$ is the ratio of natural (radiation damped) to thermal line widths

$$a \equiv \frac{\gamma}{4\pi \Delta \nu_{Dop}}.$$  

(5)

and the natural broadening width is given by the damping constant $\gamma$ which is expressed as a sum of radiative $A^r$ and autoionization $A^\alpha$ rates

$$\gamma = \sum_{l < u} A^r_{ul} + \sum_{l' < u} A^\alpha_{ul'}.$$  

(6)

$H(a, x)$ is normalized so that its integral over $x$ is $\sqrt{\pi}$.

The Voigt function must be evaluated to account for line self-shielding when finite column densities are encountered, while the total radiative decay ratio out of the upper level is needed to derive the damping constant $\gamma$. This lifetime is the sum of the autoionization and radiative decay rates. Our new data do not include calculations of autoionization rates (except for the five lines mentioned earlier). However, in order to calculate the damping constant $\gamma$ we need to assume a value, so we adopted the value, $10^{15}$ s$^{-1}$, taken from OP results for Fe xv and Fe xvi. The other data sources include autoionization rates, which we adopt.

Figure 5 shows the resulting damping parameters, plotted as a function of transition wavelength, for a gas kinetic temperature of $10^4$ K. UTA transitions are strongly damped, often with $a \gg 1$, because of the rapid autoionization rate. For comparison, the damping parameter for a strong UV line, such as H i Ly$\alpha$, is $a \sim 4 \times 10^{-4}$. It should be noted that the high damping parameter $a$ is a consequence of the inner-shell nature of these excitations regardless of their being in UTAs.

While simple approximations to the Voigt function may be sufficient for UV–optical spectroscopy, with small $a$, the large range in damping parameter that occurs when UTAs are considered make it important to use implementations of the Voigt function which are accurate for all $a$. We adopt the routine provided by Wells (1999), combined with a specially designed faster routine for $a \leq 0.1$, which provides results with a relative accuracy of $1$ in $10^4$, and has been confirmed to pass all the test cases given by Zaghloul & Ali (2011).

Stellar atmosphere texts (Rutten 2003; Mihalas 1978) often focus on results derived for approximations to the Voigt function which, while valid in the original context, are not accurate for the full range of frequency and $a$ needed to include UTA transitions. Figure 6 shows the function for a typical strong UV line such as H i Ly$\alpha$ ($a \sim 10^{-4}$) and a strongly damped UTA transition ($a \sim 10^3$). We see that at line center $H(a, x) \sim (1 + a)^{-1}$, and that the core of the line is roughly $x \sim (1 + a)$ wide.
There are a large number of lines which strongly overlap to produce the observed UTA features. The effects of mutual line shielding due to overlap must be included if the ionization rate is to be properly computed. Line overlap is treated using a combination of a coarse, low-resolution continuum, and a fine continuum, with resolution sufficient to resolve lines, as described in Shaw et al. (2005).

4. APPLICATION TO AGN

4.1. Fe xiv as a Density Indicator

Our Fe xiv data include both fine-structure levels within the ground term, namely, \( ^2 P_{1/2} \) and \( ^2 P_{3/2} \). These are separated by 2.33741 eV and produce the famous “green” Fe xiv coronal line at 5303 Å. There are approximately the same number of UTA lines originating from either level in the overall configuration.

The critical density of the excited \( J = 3/2 \) level is \( \sim 3 \times 10^9 \) cm\(^{-3} \) at 10\(^8 \) K. The temperature \( T = 10^9 \) K is approximately that of the formation of the Fe ions discussed in current paper. At considerably lower densities most of the population will be in the \( J = 1/2 \) level, while for high temperatures and densities the levels will be populated according to their statistical weight and most will be in the excited level. If the absorption characteristics are different in these two different limits then the UTA lines could be used as a density indicator.

Figure 7 (right frame) shows how the profile of the \( \lambda 15.5 \) Fe xiv UTA changes with density. The calculation assumed a total hydrogen column density of \( 10^{21} \) cm\(^{-2} \), a solar Fe abundance, and two hydrogen densities, \( n_H = 10^9 \) cm\(^{-3} \), sufficiently low for all populations to lie in the lower \( J = 1/2 \) level, and \( n_H = 10^{15} \) cm\(^{-3} \), high enough for the levels to be populated according to their statistical weights. Significant differences are present. The features originating from the configurations which include the excited \( J = 3/2 \) level are not present in the low-density profile. At the high-density limit, the ratio of the excited and lower level population is 2:1.

Therefore, new features connected to the \( J = 3/2 \) level (e.g., \( \lambda 15.35, \lambda 15.40, \) and \( \lambda 15.65 \)) become evident, whereas features originating from the ground \( J = 1/2 \) level (e.g., \( \lambda 15.30, \lambda 15.43, \) and \( \lambda 15.47 \)) become less pronounced. The total absorption does not change because the sum of the absorption oscillator strengths in this wavelength region is approximately the same both for \( J = 3/2 \) and \( J = 1/2 \) levels, although different transitions are produced by each level.

Figure 7 (left frame) shows the effects of density in the area of the \( \lambda 13.0 \) UTA lines in Fe xiv. It demonstrates very similar behavior to the case of the \( \lambda 15.5 \) UTA shown in the right panel, with \( J = 3/2 \) lines appearing at higher densities and \( J = 1/2 \) lines being prominent at low-density limit.

Future generations of X-ray spectrometers may be able to use such differences to measure the density of the absorbing gas. This would make it possible to deduce the location of the gas. Nevertheless, improved accuracy of the theoretical wavelengths would be needed for such a test to be definitive.

Absorption line spectroscopy generally has a degeneracy introduced by the fact that the ionization distribution determines an ionization parameter, the ratio of the flux of ionizing photons to gas density. Literature on this type of analysis is extensive, and summarized by Chakravorty et al. (2009). The ionization itself is not sensitive to either flux or density. By measuring the density and ionization parameter, the flux could be deduced, which then leads to the determination of the source–cloud separation.

Tests show that the density effects do not have a significant effect on the ionization of the gas because the total absorption oscillator strengths are similar. In the current implementation, we assume the low-density limit in computing the effects of UTA transitions and the resulting spectra.

The ground terms of Fe\(^{44+} \) and Fe\(^{45+} \) are simpler, having only one level, so such distinctions do not occur.

4.2. Probing the Thermal Stability of the Warm Absorber

The nature of the absorbing and emitting clouds in AGN is a long-standing problem (Osterbrock & Ferland 2006;
Figure 7. UTA spectrum of Fe\textsuperscript{xiv} transmitted through a cloud with a column density of $N(\text{H}) = 10^{21} \text{ cm}^{-2}$ and a solar Fe abundance. The effects of different densities upon the shape of the Fe\textsuperscript{xiv} UTA are shown, with the low-density case in blue having nearly all populations in the lower $J = 1/2$ level, while the high-density case has level populations within the ground term proportional to their statistical weight. This calculation only included Fe\textsuperscript{xiv}.

(A color version of this figure is available in the online journal.)

Chakravorty et al. (2009). By analogy with the local interstellar medium, several gas phases are thought to be present at one location, each in equilibrium with the radiation field, but having different levels of ionization and a range of temperatures, but the same gas pressure. This has been discussed extensively in the literature (Krolik et al. 1981; Hess et al. 1997; Krolik & Kriss 2001; Komossa & Mathur 2001; Reynolds & Fabian 1995).

The ionization, temperature, and spectrum of clouds that can exist is determined by the type of stability analysis shown in Figure 8. This shows the familiar “S curve,” which is computed for an optically thin cell of gas exposed to the AGN radiation field. Thermally stable phases have positive slope, while regions with negative slope are unstable so that gas will only exist in these regions for a short time. The shape of the S curve determines the properties of clouds which are thermally stable and so live long enough to contribute to the observed spectrum.

Recent S-curve calculations have focused on how properties of the AGN might affect its shape, and so determine which clouds might exist. The literature on this topic is vast, and recent examples include Holczer et al. (2007, 2010) and Holczer & Behar (2012), who examine observational determinations of the stable points on the S curve. Chakravorty et al. (2008), who show the effects of updated dielectronic recombination rate coefficients, Chakravorty et al. (2009), who do a systematic stability analysis, and Chakravorty et al. (2012), who discuss the influence of the spectral energy distribution (SED) on the cloud stability.

Here, we show the effects of the improvements discussed in this paper, using version C13 of Cloudy (Ferland et al. 2013). Some other recent improvements to the atomic data, which focus on the atomic models used to compute the cooling, are described by Lykins et al. (2013). Figure 8 shows the thermal stability curve resulting from the improved atomic physics in this paper. Many model parameters are taken from Chakravorty et al. (2009). The $x$-axis is the ratio of the dimensionless ionization parameter $U$, defined as the ratio of ionizing photon to hydrogen densities, to the gas kinetic temperature, while the latter is the $y$-axis. Thermally stable regions, those with positive slope, are shown as the heavier lines.

In a series of papers Holczer et al. (2007, 2010), and Holczer & Behar (2012) infer from observations of column densities of certain ions that temperatures between $4.5 < \log T < 5.0$ are missing and ascribe it to thermal instability. On the other hand,
Figure 9. Fe ionization distribution across Figure 8, where the gas kinetic temperature is indicated by the solid line increasing from left to right. Horizontal lines at the top of the figure mark the thermally stable regions of Figure 8. The Fe ions considered in this paper probe the high temperature end of the cool branch in Figure 8, unstable regions, and the low temperature end of the middle stable branch.

(A color version of this figure is available in the online journal.)

Figure 10. Ratio of the UTA ionization to total ionization rate for several Fe charge stages. Filled circles include the results presented in this paper, while the crosses represent the older data. These are for the conditions occurring across Figure 8. Rates are evaluated at the illuminated face so line self-shielding is not important. UTA ionization is the dominant ionization process for the ions considered in this paper.

(A color version of this figure is available in the online journal.)

Figure 11. Spectrum of a cloud lying along the line of sight to an AGN. Cloud parameters are indicated in the upper panel, which shows the transmitted coarse continuum. The lower panel shows a small portion of the fine continuum with predictions using the atomic data summarized in this paper in black, and with older UTA line data in green.

(A color version of this figure is available in the online journal.)

this region is stable in the current calculation with its assumed parameters. The shape of the stability curve is affected by several other ingredients besides the atomic data. The composition assumed, and the form of the SED, also change it. This suggests that the presence or absence of stable gas could be used to infer the SED or gas composition, among other properties of the AGN.

Figure 9 shows the distribution of ionization stages of Fe as a function of the ionization parameter $U$. The ions discussed in this paper peak in the range $\log U \sim 1−2$. These probe the upper bound of the low-$T$ branch of thermally stable gas, the lowest unstable region, and the low-$T$ part of the middle stable branch. Horizontal lines in the upper part of Figure 9 indicate the regions where the gas is stable.

The atomic data presented in this paper affect the details of the transition between the cool and middle stable branches. Figure 10 shows the ratio of UTA to total ionization for several charge states of Fe, where the filled circles include the data presented in this paper, while the crosses represent the older data. The larger number of lines derived here results in a larger UTA ionization rate, as shown in the figure. By UTA ionization we mean the effect of autoionization following inner-shell photoexcitation. UTAs have the greatest effect on the ions which occur around the transition from the cool to middle stable branch, and their physics affects the details of this transition.

Finally, in Figure 11 we show the spectrum produced by an intervening cloud near the upper range of the cool stable branch. This cloud has solar abundances, a column density of $N(H) = 10^{21}$ cm$^{-2}$, and an ionization parameter of $\log U = 1.25$. A portion of the coarse continuum which is used for continuum radiative transfer and atomic rates is shown in the upper panel. The assumed SED, which includes the “Big Bump” from the central accretion disk and a non-thermal X-ray power law, has many absorption features superimposed. Emission lines are also produced by the cloud but are weak in this portion of the spectrum due to the low gas temperature.

The lower panel shows a small portion of the fine continuum in the neighborhood of the $\sim 15$ Å UTA feature. Two curves are shown, with the solid line using the results presented in this
paper, and the dashed line employing the other data sources summarized above. Significant differences are present.

5. CONCLUSIONS

This paper has summarized advances in the atomic data needed to simulate conditions in a photoionized plasma. All of these improvements are now in the development version of the plasma simulation code Cloudy, and will be part of the next major release. The specific advances are the following.

1. We have calculated a large set of atomic data for UTA lines using the CI method implemented in the CIV3 code. This new data set substantially extends previous atomic data of Kisielius et al. (2003) in two ways. First, the E1 transitions from the inner 2s shell are determined. Second, the data sets for Fe xiv, Fe xv, and Fe xvi include lines 2l – 4l, 5l in addition to the earlier determined lines 2p – 3l.1.

2. We have incorporated this large set of new UTA data into the simulation code Cloudy, and applied it to problems in AGN. These improvements will be part of the next major release of the code.

3. We summarize our data sources for UTA transitions. There are still missing data, even for very important ions. These should be a priority for new theoretical calculations.

4. We summarize how our data, which were computationally very expensive to undertake, compare to simpler calculations. Line wavelengths differ (although insignificantly) due to differences in the computed level energies, but the transition rates are in good agreement.

5. The UTA lines are often strongly damped, many having damping parameters a ≫ 1. We have improved the form of the Voigt function used by Cloudy to handle such strongly damped lines.

6. We show how the Fe xiv UTA at λ15.5 can be used to measure the density of the gas, or identify whether the density is significantly above or below 10⁹ cm⁻³. Such measurements would help determine the location of the warm absorber in AGN.

7. The total ionization rate is increased by roughly 30% with the new set of UTA data, which have far more lines. This changes the ionization of the gas and alters the thermal properties of photoionized gas exposed to a typical AGN SED.

8. We present a newly computed thermal stability “S curve” using the new data. We show that the Fe ions considered in this paper are produced in the warmer parts of the cool thermally stable branch, an unstable region, and in cooler parts of the middle stable branch. As a result, these lines probe the portions of the S curve which determine which cloud parameters can persist. Future work will investigate the effects changes in these regions have upon the observed spectrum.

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