Mg Line formation in late-type stellar atmospheres
I. The model atom

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ABSTRACT

Context. Magnesium is an element of significant astrophysical importance, often traced in late-type stars using lines of neutral magnesium, which is expected to be subject to departures from local thermodynamic equilibrium (LTE). The importance of Mg, together with the unique range of spectral features in late-type stars probing different parts of the atom, as well as its relative simplicity from an atomic physics point of view, makes it a prime target and test bed for detailed ab initio non-LTE modelling in stellar atmospheres. Previous non-LTE modelling of spectral line formation has, however, been subject to uncertainties due to lack of accurate data for inelastic collisions with electrons and hydrogen atoms.

Aims. In this paper we build and test a Mg model atom for spectral line formation in late-type stars with new or recent inelastic collision data and no associated free parameters. We aim to reduce these uncertainties and thereby improve the accuracy of Mg non-LTE modelling in late-type stars.

Methods. For the low-lying states of Mg I, electron collision data were calculated using the R-matrix method. Hydrogen collision data, including charge transfer processes, were taken from recent calculations by some of us. Calculations for collisional broadening by neutral hydrogen were also performed where data were missing. These calculations, together with data from the literature, were used to build a model atom. This model was then employed in the context of standard non-LTE modelling in 1D (including average 3D) model atmospheres in a small set of stellar atmosphere models. First, the modelling was tested by comparisons with observed spectra of benchmark stars with well-known parameters. Second, the spectral line behaviour and uncertainties were explored by extensive experiments in which sets of collisional data were changed or removed.

Results. The modelled spectra agree well with observed spectra from benchmark stars, showing much better agreement with line profile shapes than with LTE modelling. The line-to-line scatter in the derived abundances shows some improvements compared to LTE (where the cores of strong lines must often be ignored), particularly when coupled with averaged 3D models. The observed Mg emission features at 7 and 12 μm in the spectra of the Sun and Arcturus, which are sensitive to the collision data, are reasonably well reproduced. Charge transfer with H is generally important as a thermalising mechanism in dwarfs, but less so in giants. Excitation due to collisions with H is found to be quite important in both giants and dwarfs. The R-matrix calculations for electron collisions also lead to significant differences compared to when approximate formulas are employed. The modelling predicts non-LTE abundance corrections Δ(Mg)NLTE in dwarfs, both solar metallicity and metal-poor, to be very small (of order 0.01 dex), even smaller than found in previous studies. In giants, corrections vary greatly between lines, but can be as large as 0.4 dex.

Conclusions. Our results emphasise the need for accurate data of Mg collisions with both electrons and H atoms for precise non-LTE predictions of stellar spectra, but demonstrate that such data can be calculated and that ab initio non-LTE modelling without resort to free parameters is possible. In contrast to Li and Na, where only the introduction of charge transfer processes has led to differences with respect to earlier non-LTE modelling, the more complex case of Mg finds changes due to improvements in the data for collisional excitation by electrons and hydrogen atoms, as well as due to the charge transfer processes. Grids of departure coefficients and abundance corrections for a range of stellar parameters are planned for a forthcoming paper.

Key words. atomic data – atomic processes – line: formation – stars: abundances – stars: late-type

1. Introduction

Neutral magnesium creates a broad range of spectral features in late-type stars, including some of the strongest lines in their spectra. Consequently, Mg is detectable even in low-quality spectra and in metal-poor stars, making Mg an excellent tracer of α-element abundances. The strong Mg I λ lines with pressure-broadened wings can be used as a surface gravity diagnostic (e.g. Edvardsson 1988; Fuhrmann et al. 1997). Additionally, the inter-combination line at 4571 Å is unique among lines of similar strength in the Sun in that it is expected to have an almost LTE line formation across the entire profile (see Carlsson et al. 1992, and discussion therein). Emission features due to Mg I Rydberg transitions have been identified in the infrared (IR) region of the solar spectrum, particularly those near 12 μm (Murcray et al. 1981; Chang & Noyes 1983). The Mg I emission

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features dominate other features arising from similar transitions in other atoms due to a combination of abundance and ionisation energy. These emission lines have also been observed in other main-sequence and giant stars: Arcturus (Uitenbroek & Noyes 1996), Procyon (Ryde et al. 2004), and Pollux (Sundqvist et al. 2008).^1^ At \( T \geq 5000 \) K, Mg I is a minority species and as a result is expected to be sensitive to departures from LTE in stellar atmospheres, in particular due to photoionisation by non-local radiation. While in the Sun non-LTE effects on lines in the optical range are expected to be relatively weak (Zhao et al. 1998), significant departures from LTE have been predicted in metal-poor dwarfs and giants due to the increase in UV radiation because of the decreased line-blanketing (Zhao & Gehren 2000). Moreover, dwarfs and giants due to the increase in UV radiation because of significant departures from LTE have been predicted in metal-poor atmospheres, in particular due to photoionisation by non-local radiation. Furthermore, changes of the Milky Way. The ongoing Gaia satellite, several complementary ground-based spectroscopic surveys have been initiated. Their goal is to determine high-precision abundances for eventually hundreds of thousands of stars and thereby unravel the formation and evolution of the Milky Way. The ongoing Gaia-ESO Survey will determine [Mg/Fe] ratios in all Galactic components to better than \(-0.1\) dex, in order to distinguish between the thin and thick disk and between accreted and in-situ halo components, for instance (Recio-Blanco 2013; Gilmore 2012). The targeted wavelength regions of the VLT/GIRAFFE spectrograph contain two strong Mg lines, 5528 Å and 8806 Å, detectable in all FGK type stars down to low metallicities. For stars in the solar neighbourhood, the GALAH survey (De Silva et al. 2015) aims at a higher precision (\(-0.05\) dex) for many elements with the aim to identify dispersed star clusters (Zucker et al. 2012). The wavelength settings of the AAT/HERMES instrument contain two weak Mg lines, 5711 Å and 7691 Å, suitable for analysis of high-metallicity stars.

^1^ The 12.3 \( \mu m \) feature is observed in absorption in Betelgeuse (Uitenbroek & Noyes 1996), but this is due to the blending with a strong water line (Sundqvist et al. 2008).

2. Previous works

As mentioned, a significant number of non-LTE Mg I line formation studies have been conducted, and the main difference between these, especially the more recent and comprehensive studies since around 1990, and the work presented here lies with the atomic collision data for bound-bound processes. Thus, as a basis for comparison, in this section we present an overview of some past studies, focussing on the atomic collisional data employed.

The earliest investigation of Mg I line formation by Athay & Canfield (1969) examined the Mg b lines in the Sun. They used a small model atom with constant values for the electron collisional cross sections based on the van Regemorter (vR) formula (van Regemorter 1962) – an often-used semi-empirical interpolation formula based on the Bethe approximation. Altrock & Canfield (1974) updated the collisional cross-sections to those calculated by van Blerkom (1970). Mauas et al. (1988) used a 13-level model atom to study the 4571 Å line and the Mg b line at 5173 Å in the quiet Sun employing updated electron collisional data collected from various experiments and calculations.

Subsequent studies in the late 1980s and early 1990s focussed on the Mg I IR emission features at 12 and 18 \( \mu m \) observed in the Sun (Lemke & Holweger 1987; Chang et al. 1991; Carlsson et al. 1992, among others), where high-lying and Rydberg levels were included. Chang et al. used a model atom with 41 levels where 3s \( nl \) levels with \( l = s, p, d \) and \( n \leq 7 \) are included, and 7f, 7g, 7h, 7i are merged; \( 8 \leq n \leq 15 \) are super levels. Carlsson et al. presented a model atom with 71 levels plus continuum, including up to 3s \( nl \) with \( n \leq 9, l \leq 8 \), while for \( n = 10, l = s, p \) levels are included separately and \( l \geq 2 \) are included as single super levels. For \( l \geq 3 \) singlet and triplet terms are collapsed to single levels. These studies demonstrated the importance of comprehensive models including levels up to the quasi-continuum, where levels are collisionally dominated and strongly coupled to the continuum. This permits a population flow from the continuum to lower-lying levels. The importance of this was demonstrated by the failure of Lemke & Holweger’s model to reproduce the 12 \( \mu m \) emission, due to a lack of levels sufficiently close to the continuum to be collisionally dominated, in addition to incorrect collision rates (see Sect. 6.6 of Carlsson et al.). Sundqvist et al. (2008) also showed the need to extend the model atoms to include \( n \geq 10 \) for giant stars; the sparser atmosphere means that the levels become collisionally dominated at higher \( n \).

Chang et al. and Carlsson et al. both took electron collisional excitation rates from Mauas et al. For the optically allowed transitions not included in Mauas et al., Chang et al. used the vR formula, while Carlsson et al. used the impact parameter (IP) method (Seaton 1962). For forbidden transitions, Chang et al. used the vR formula with an arbitrary oscillator strength,
while Carlsson et al. used an arbitrary scaling of nearby allowed transitions.

Inelastic collisions with neutral hydrogen atoms were included in Chang et al. (1991). For Mg\textsc{i}+H collisional excitation the formula from Kaulakys (1985) was used for transitions between Rydberg levels, but did not include data for the lower-lying Mg\textsc{i} levels. Charge-transfer processes involving Rydberg states and protons, Mg\textsc{ii}(n\ell)+H\rightarrow Mg\textsc{iii}+H(n), are disregarded, but suggested to be unimportant in the solar photosphere, despite the large rate coefficients, due to low number densities. We note that we have been unable to find the source of these data quoted as being in preparation. Carlsson et al. discussed the possible role of collisional excitation with neutral H, arguing that they will lead to significant \( l \)-changing collisions among Rydberg states. They included this effect by setting high electron collision rates between levels of equal \( n \) and with small differences in quantum defects to ensure relative LTE between these levels. Various revisions of the model atom in Carlsson et al. have been used for studying the 12 and 18 \( \mu \)m features in stars other than the Sun (e.g. Ryde et al. 2004; Sundqvist et al. 2008), some including collisions with hydrogen atoms via the Drawin formula (Drawin 1969; Steenbock & Holweger 1984). The Drawin formula has been shown in recent years to be unable to provide reliable estimates of hydrogen collision processes, and no estimates at all for optically forbidden transitions or charge transfer.

Various revisions of the model atom in Carlsson et al. have been used for studying the 12 and 18 \( \mu \)m lines and demonstrated that the shapes of these lines are relatively insensitive to granulation-induced asymmetries. That work is, however, to our knowledge the only attempt at 3D non-LTE Mg line formation to date. We further note that Rutten & Carlsson (1994) provided an excellent summary of non-LTE mechanisms in Mg\textsc{i}, and Chang (1994) provided a summary of the corresponding atomic physics relating to the infrared lines.

3. Non-LTE modelling

We performed standard 1D non-LTE modelling, where the coupled radiative transfer and statistical equilibrium equations based on our Mg model atom are solved in 1D stellar model atmospheres using version 2.3 of the MULTI code (Carlsson 1986, 1992). We employed the same background line opacity data (Collet et al. 2005) used in the calculation of the MARCS theoretical stellar atmospheres (Gustafsson et al. 2008), but resampled to 10 300 frequency points. Turbulence was modelled by the usual micro- and macro-turbulence parameters.

The MULTI code treats the restricted non-LTE radiative transfer problem, in which the element of interest, here Mg, is treated as a trace element. Thus, it is assumed that there is no feedback on the atmospheric structure or background opacities, which are computed assuming LTE. Only very few studies have treated the full non-LTE problem in cool stars (Short & Hauschildt 2005, 2009, Lind et al. 2012) considered the effects of increased electron densities and decreased UV opacity due to the increased ionisation of Fe\textsc{i} on the non-LTE line formation of Fe, arguing that the first effect was insignificant and the second weaker than typical non-LTE corrections. However, such tests are limited, and as stated by the authors, only fully consistent calculations will resolve this definitively. Such calculations are beyond the authors’ present capabilities and beyond the scope of the present work. We note that the abundance of Mg is almost always larger than Fe, especially in metal-poor stars (typically about a factor of 3), and Mg also has a somewhat lower ionisation energy, making it a more important electron donor. On the other hand, non-LTE does not perturb the ionisation balance of Mg as much as for Fe. It would be important to test the trace element assumption.

In this section we present the details of our atomic model, often referred to as the “model atom”. In Sect. 3.1 we discuss the energy levels and the size of the model atom. In Sect. 3.2 the radiative data are outlined, namely bound-bound (b-b) and bound-free (b-f) transition probabilities and line broadening cross sections, and in Sect. 3.3 the inelastic collision data are described. In Sect. 3.4 we describe the modified model atoms used to test the influence of new data on the final results.

3.1. Energy levels

Following earlier studies, we decided to build a model atom including Mg\textsc{i}, Mg\textsc{ii}, and the ground state of Mg\textsc{iii}. To build the structure of the model atom, energy level data were collected from the NIST database (Ralchenko et al. 2010), supplemented with data for high \( n \) (up to \( n = 80 \)) in the case of Mg\textsc{i} from Kurucz & Bell (1995). For high angular momentum levels absent from either the NIST or Kurucz & Bell collections, we made use

\[ \text{Macroturbulence is assumed to be Gaussian and treated according to the radial-tangential model.} \]
of the Sommerfeld-Dirac expression to estimate the energies. In principle, it is desirable to have the model atom as large and as detailed as possible; however, this is neither computationally practical nor required to achieve converged results for the spectral lines of interest. Thus, to investigate the convergence of the model atom, a computer script was written to generate model atoms characterised by three parameters: 1) the largest principle quantum number for which levels are included, 2) the principle quantum number above which all levels are merged into “super levels” encompassing all states of same \( n \), and 3) the term above which all fine-structure components \( J \) are merged into a single level. This permitted to test a range of atomic models in size and detail.

Finally, after extensive testing, we found that the following model atom provided results that converged to better than 1% for a range of late-type stellar parameters. \( \text{Mg} \text{I} \) includes states up to \( n = 20 \) and uses super levels for \( n > 10 \). Fine-structure splitting was implemented for \( ^3P \) states up to \( 3s \ ^3P_{0,1,2} \), as well as for the \( 3p^2 \ ^3P_{0,1,2} \) states. Splitting was not included for states \( 3s \ ^1L \ ^3L \) where \( L \geq 2 \). \( \text{Mg} \text{II} \) includes states up to \( n = 11 \) and uses super levels for \( n > 7 \). For states up to \( 4f^2\ ^2F_{5/2,7/2} \) fine-structure splitting was included. This gives a final model atom covering \( \text{Mg} \text{I} \), \( \text{Mg} \text{II} \), with 108 and 34 states respectively, plus the ground state of \( \text{Mg} \text{III} \). Grotrian diagrams for \( \text{Mg} \text{I} \) and \( \text{Mg} \text{II} \) are shown in Fig. 1.

### 3.2. Radiative data

Radiative b-b transition probabilities were taken mainly from NIST and Kurucz & Bell (1995). When data for a transition were in both sources, the data from NIST were adopted. For the \( \text{Mg} \text{II} \) singlet, oscillator strengths were taken from Aldenius et al. (2007). When transitions involving Rydberg states of \( \text{Mg} \text{I} \) were needed and data were unavailable, transition probabilities were calculated following Guseinov & Mamedov (2012). The data for the most important diagnostic lines are provided in Table 1. The final model atom has 1185 b-b transitions (962 \( \text{Mg} \text{I} \) and 223 \( \text{Mg} \text{II} \)). In all cases, radiative damping was calculated consistently from the employed transition probabilities.

For most UV lines, \( S_n > B_n \), in the line formation region contributing to UV overionisation that tends to depopulate lower levels. Thus, b-f processes are of considerable importance. Bound-free cross-sections for \( \text{Mg} \text{I} \) and \( \text{Mg} \text{II} \) were taken from TOPbase (Cunto & Mendoza 1992; Cunto et al. 1993). The data for \( \text{Mg} \text{I} \) were adopted from Butler et al. (1993), and are available for \( n \leq 9 \) and \( l \leq 4 \). For \( \text{Mg} \text{II} \) the original source is unpublished and data are available for \( n \leq 10 \) and \( l \leq 3 \). For larger \( n \) and \( l \), hydrogenic cross-sections were used. The threshold energies of the photoionisation cross-sections taken from TOPbase do not coincide exactly with the experimental ones used in our modelling, and so the TOPbase cross-sections were shifted appropriately. Strong resonances in the b-f cross sections sometimes play an important role. The photoionisation cross-sections for the ground and first excited state of \( \text{Mg} \text{I} \) have no resonances strong enough to be important at typical temperatures in cool stellar atmospheres. The photoionisation from the \( 3s3p \ ^1P \) level shows a resonance at \( \lambda \sim 2900 \AA \) due to the \( 3p^2 \ ^1S \) auto-ionising level that is five times greater than the value of the cross-section at threshold, thus making it an important contributor. The photoionisation from the \( 3s4s \ ^1S \) levels shows resonances due to the \( 3p4s \ ^1P \) auto-ionising levels that are more than 500 times greater than the value at threshold, making these resonances the main contributors to the total photoionisation rate at temperatures around 5000 K. \( \text{Mg} \text{II} \) photoionisation cross-sections do not have resonances in the energy range of interest.

#### 3.2.1. Line broadening

In addition to turbulent and radiative broadening, both mentioned above, data on the effects of collisions on the modelled spectral lines were also required, in particular broadening of the lines due to collisions with charged particles and neutral hydrogen atoms. For lines of diagnostic interest in this work (see Table 1), these data need to be as accurate as possible, while for other transitions this is probably not of great importance, and reasonable estimates suffice.

For some of the \( \text{Mg} \text{I} \) lines we included collisional broadening due to electrons and protons using quadratic Stark broadening parameters from Dimitrijevic & Sahal-Bréchot (1996). We adopted the line width at 5000 K neglecting variation with temperature, which is typically weak. For the 18 \( \mu \)m lines we adopted widths from van Regemorter & Hoang Binh (1993). For the inter-combination line \( 3p^2 \ ^3P \sim 3s \ ^1S \) at 4571 Å we adopted the value used in Mashonkina (2013)\(^3\). For all other lines (i.e. those not in Table 1, or of \( \text{Mg} \text{II} \), Stark broadening was neglected.

For transitions of \( \text{Mg} \text{I} \) involving low-lying states, collisional broadening due to collisions with neutral H is described via cross-sections and velocity parameters interpolated in tables.

\(^3\) This value is stated to be taken from the VALD database, but we were unable to find any such data for this line in the VALD.
Table 1. Data for important transitions, including diagnostic lines.

| \( \lambda \) | Transition | \( \log gf \) | Source | \( \sigma \) | \( \alpha \) | \( \log \Gamma_\alpha/N_\alpha \) (5000 K) | Source | Dr | Dp | Gr | Gp |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| [Å] | | | | | | [rad s\(^{-1}\) cm\(^{3}\)] | | | | | |
| Mg I | | | | | | | | | | | | |
| 3829 | 3d \( ^{1}D \rightarrow 3p \, ^{1}P \) \(_{0} \) | 0.23 | Fro/NIST | 708 | 0.301 | 4.51 | D-SB | H | H | CH | CH |
| 3832 | 3d \( ^{1}D \rightarrow 3p \, ^{1}P \) \(_{0} \) | 0.52 | Fro/NIST | 708 | 0.301 | 4.51 | D-SB | H | H | CH | CH |
| 3838 | 3d \( ^{1}D \rightarrow 3p \, ^{1}P \) \(_{2} \) | 0.47 | Fro/NIST | 708 | 0.301 | 4.51 | D-SB | H | H | CH | CH |
| 4167 | 7d \( ^{3}D \rightarrow 3p \, ^{1}P \) | 0.75 | C-T/NIST | 222 | 0.249 | 3.49 | D-SB | H | H | CH | H |
| 4571 | 3p \( ^{3}P \rightarrow 3s \) \(_{1} \) | 5.62 | Fro/NIST | 222 | 0.241 | 5.51 | H | H | CH | CH |
| 4703 | 5d \( ^{1}D \rightarrow 3p \, ^{1}P \) | 0.44 | C-T/NIST | 2806 | 0.269 | 4.11 | D-SB | H | H | CH | H |
| 5167 | 4s \( ^{3}S \rightarrow 3p \, ^{1}P \) \(_{0} \) | 0.93 | Ald | 728 | 0.238 | 5.40 | D-SB | H | CH | CH | CH |
| 5173 | 4s \( ^{3}S \rightarrow 3p \, ^{1}P \) \(_{1} \) | 0.45 | Ald | 728 | 0.238 | 5.40 | D-SB | H | CH | CH | CH |
| 5184 | 4s \( ^{3}S \rightarrow 3p \, ^{1}P \) \(_{2} \) | 0.24 | Ald | 728 | 0.238 | 5.40 | D-SB | H | CH | CH | CH |
| 5528 | 4d \( ^{1}D \rightarrow 3p \, ^{1}P \) | 0.50 | C-T/NIST | 1460 | 0.312 | 4.56 | D-SB | H | H | CH | CH |
| 8710 | 7d \( ^{3}D \rightarrow 4p \, ^{3}P \) \(_{0} \) | 1.57 | But/NIST | 2.72 | D-SB | CH | H | CH | H |
| 8711 | 7d \( ^{3}D \rightarrow 4p \, ^{3}P \) \(_{1} \) | 1.09 | But/NIST | 2.72 | D-SB | CH | H | CH | H |
| 8718 | 7d \( ^{3}D \rightarrow 4p \, ^{3}P \) \(_{2} \) | 0.87 | But/NIST | 2.72 | D-SB | CH | H | CH | H |
| 8736 | 3d \( ^{3}D \rightarrow 3d \, ^{3}D \) | 0.53 | But/NIST | 2.95 | D-SB | CH | H | CH | H |
| 8806 | 3d \( ^{3}D \rightarrow 3p \, ^{1}P \) | 0.13 | Fro/NIST | 529 | 0.277 | 5.39 | D-SB | CH | CH | CH | CH |
| 7370 | 6h \( ^{2}H \rightarrow 5g \, ^{2}G \) | 1.34 | Civ | 4950 | 1.549 | 3.06 | vR-HB | CH | CE | CH | CH |
| 12270 | 7h \( ^{2}H \rightarrow 6g \, ^{2}G \) | 1.62 | Civ | 5191 | 1.738* | 2.39 | D-SB | CH | CH | CH | CH |
| 12320 | 7i \( ^{1}H \rightarrow 6h \, ^{2}H \) | 1.95 | Hydro | 4657 | 1.752* | 2.55 | D-SB | H | CH | CH | CH |
| 188300 | 8h \( ^{2}H \rightarrow 7g \, ^{2}G \) | 0.49 | Hydro | 4497 | 1.764* | 2.92 | vR-HB | CH | H | CH | CH |
| 189500 | 8i \( ^{1}H \rightarrow 7h \, ^{2}H \) | 1.90 | Hydro | 4304 | 1.778* | 2.12 | vR-HB | H | CH | CH | CH |

Mg II

| \( \lambda \) | Transition | \( \log gf \) | Source | \( \sigma \) | \( \alpha \) | \( \log \Gamma_\alpha/N_\alpha \) (5000 K) | Source | Dr | Dp | Gr | Gp |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| | | | | | | | | | | | | |
| 4385 | 5d \( ^{3}P \rightarrow 4p \, ^{3}P \) | 0.78 | Sie/NIST | 1 | 0.29 | Sie/NIST |
| 4391 | 3d \( ^{3}D \rightarrow 4p \, ^{3}P \) | 0.48 | Sie/NIST | 1 | 0.29 | Sie/NIST |
| 4481 | 4f \( ^{3}P \rightarrow 3d \, ^{3}D \) | 0.76 | Fro/NIST | 1 | 0.29 | Sie/NIST |
| 5402 | 7g \( ^{3}G \rightarrow 4f \, ^{3}F \) | 0.06 | Mas | 1 | 0.29 | Sie/NIST |
| 7877 | 4d \( ^{3}D \rightarrow 4p \, ^{3}P \) | 0.39 | Sie/NIST | 1 | 0.29 | Sie/NIST |
| 7896 | 4d \( ^{3}D \rightarrow 4p \, ^{3}P \) | 0.64 | Sie/NIST | 1 | 0.29 | Sie/NIST |

Notes. Columns 1 and 2 give the transition wavelength and labels for the states involved. Columns 3 and 4 give the adopted oscillator strength, in the form \( \log gf \), and the source of these data (see notes). Columns 5–8 give collisional broadening data (see text and notes). Columns 9–12 show, for each line of Mg I, the collision process most responsible for the changes between model B and model F, and thus indicates the most important contributor of the new collision data introduced in this work (see Sect. 5 for more details). Results are given for the four test atmospheric models described in Sect. 5: Dr = Dwarf, metal-rich; Dp = Dwarf, metal-poor; Gr = Giant, metal-rich; Gp = Giant, metal-poor. The collisional processes are labelled (see Sect. 5) H when the line has a sensitivity similar to charge transfer with H and CH; collisional excitation with H, CH, and CE indicates the line is sensitive mostly to the electron collision rates. For Mg II lines there are non-LTE effects in some lines, but the non-LTE abundance corrections are not sensitive to any particular collisional process. Oscillator strengths (\( f \)-values) were calculated mostly from the NIST database (Ralchenko et al. 2010). The original sources of the NIST data are Froese Fischer & Tachiev (2010) [Fro], Chang & Tang (1990) [C-T] Butler et al. (1993) [But], Siegel et al. (1998) [Sie]. Other sources are Aldenius et al. (2007) [Ald], Civiš, S. et al. (2013) [Civ] and Kurucz & Peytremann (1975) [K-P]. For the Mg I IR lines with no data found in the literature we calculated the \( f \)-values using the hydrogenic formula from Guseinov & Mamedov (2012) [Hydro]. To calculate the van der Waals line widths \( \Gamma_i \) we used data in the ABO theory format where \( \sigma \) is the broadening cross-section in atomic units and \( \alpha \) is the velocity parameter. These were calculated with the ABO theory except for those marked with an asterisk, which were calculated in this work (see Table 3 for the IR lines). Stark broadening line widths, \( \Gamma_i \), were taken from Dimitrijevic & Sahal-Bréchot (1996) [D-SB]; van Regemorter & Hoang Binh (1993) [vR-HB] and Mashonkina (2013, including only electrons) [Mas].
Table 2. Inelastic $l$-changing ($\sigma^l_\ell \sigma$) and $n$-changing ($\sigma^n_\ell \sigma$) cross-sections for levels of interest, in atomic units.

| Level | $\sigma^l_\ell \sigma$ [au] | $\sigma^n_\ell \sigma$ [au] |
|-------|-----------------|-----------------|
| 5g    | 4752.0          | 78              |
| 6g    | 5456.0          | 130             |
| 6h    | 4938.0          | 132             |
| 7g    | 4744.0          | 196             |
| 7h    | 4600.0          | 196             |
| 7i    | 4048.0          | 196             |
| 8h    | 3786.0          | 270             |
| 8i    | 3542.0          | 270             |
| 8k    | 3146.0          | 270             |

Notes. Columns 2 and 3 give results from this work at $v = 10^4$ m/s. For direct comparison with $l$-changing cross-sections from Table 4 of HBvR, results are also given for two states at $v = 10.457$ m/s corresponding to the average collision velocity at $T = 5000$ K. Column 4 gives our results and Col. 5 the results from HBvR.

and thus is simply the sum of the two contributions. The elastic scattering involving the two levels, however, subtracts coherently and must be treated together at the scattering amplitude level. The inelastic cross-sections may for convenience be split into $l$-changing and $n$-changing collisions, such that

$$\sigma^\text{inel}_{nl} = \sum_{\ell,ii} \sigma^\ell_{nl-\text{tot}} = \sum_{\ell,ii} \sigma^\ell_{nl-\text{tot}} + \sum_{\ell,nn} \sum_{ii} \sigma^\ell_{nl-\text{tot}}. \quad (2)$$

The cross-sections for levels in transitions of interest were calculated and are presented in Table 2. The $l$-changing inelastic cross-sections were calculated using the method for inelastic Rydberg-hydrogen collisions due to Kaulakys (1991) described in Sect. 3.3.2. The $n$-changing inelastic cross-sections were calculated in the scattering length approximation using the analytic expressions of Lebedev & Marchenko (1987, Eq. (25)) and Kaulakys (1985, Eqs. (4), (5)), which give practically identical results, as pointed out by Lebedev and Marchenko. We could calculate the $n$-changing cross-sections in the same way as done for the $l$-changing; however, the computational time is significant, and as seen in Table 2, the $n$-changing contribution is very weak, as expected (e.g. Hoang Binh & van Regemorter 1995). HBvR showed that the ratio of the elastic to the inelastic component is expected to be of order $\Delta n/n^2$, and thus we did not include the elastic contribution.

The total broadening cross-sections are then calculated according to Eq. (1) and are presented in Table 3. The cross-sections for a collision velocity of $10^4$ m/s are presented, together with a velocity parameter $\alpha$ that was calculated by fitting a power-law behaviour of the cross-section with velocity following Anstee & O’Mara (1995). Line widths are also presented and compared with the results of HBvR for the 12 $\mu$m lines. As seen from Table 2, our cross-sections are generally larger than those of HBvR, although not quite as large as their calculations using the method of Kaulakys (1991). Note that we have included the $n$-changing component, which was neglected by HBvR. Our cross-sections for the 12.2 and 12.3 $\mu$m lines differ from theirs by 1 and 21 per cent, respectively. The line widths differ by considerably more because HBvR used an approximate method for calculating the line widths from a cross-section at a single velocity, rather than an integration over a Maxwellian velocity distribution. In any case, we conclude that the uncertainty in the broadening cross-sections for these lines involving Rydberg states is at least 20 per cent, probably larger due to approximations inherent to both methods.

For transitions covered neither by the ABO theory nor by the calculations for infrared lines described above, the Unsöld formula was used with an enhancement factor of 2.5. This includes all transitions of Mg II.

3.3. Inelastic collision data

In this subsection we describe the data adopted for inelastic collision processes: collisional excitation and ionisation due to electrons, and collisional excitation and charge transfer processes due to hydrogen atoms. This includes the description of new calculations for collisional excitation of Mg I using the R-matrix method.

Figure 5 visualises the total downward collision rates calculated in the $<3D_0>_{1D}$ atmospheric model (see Sect. 4.1) at a depth corresponding to roughly $T = 6000$ K ($\log \tau_{\text{tot}} \sim -0.23$) using our final model atom. The collision matrix for Mg I has a structure very similar to that seen in Carlsson et al. (1992). In particular, we see the block structure of the very strongest transitions along the diagonal caused by the strong $l$-changing collisions where $\Delta n = 0$, where collisions with $\Delta l = 1$ are strongest of all. We also see structure off the diagonal due to the strength of collisions with $\Delta l = 1$ and $\Delta n = 1$. A new feature in this work is the structure seen in the collisional coupling of Mg I states to the Mg II ground state. Inclusion of charge transfer processes leads to rates between the levels around $4s \, ^1\text{S}$ and the Mg II ground state having the same order of magnitude as the collisional rates from these levels to neighbouring levels.

Before we describe the data, a word on the $3p^3 \, ^3\text{P}$ level is warranted. This is the only level included in our modelling where the dominant configuration is not $3s \, n\ell$. This level is too highly excited to be included in the detailed quantum mechanical calculations, and it is questionable whether the approximate methods are applicable to transitions involving two electrons. Our expectation is that collisional processes involving this state will be relatively weak, and thus we ignored collisional coupling to this state (as can be seen in Fig. 5). This causes this level to behave quite differently from other levels of Mg I in the non-LTE modelling and to depart from equilibrium very deep in the atmosphere ($\log \tau_{\text{tot}} \sim 1$ in the solar case). However, this is not important for any lines observed in stellar spectra.

3.3.1. Electrons

Collisional excitation and de-excitation between low-lying states of Mg I are accounted for with $R$-matrix calculations, which are presented in detail below. These calculations provide data for the transitions between the lowest ten states of Mg I, up to $5p \, ^1\text{P}$. For transitions involving higher lying states, including Rydberg levels, we must use more approximate methods. For allowed transitions, two such methods are in common use in stellar astrophysics: the vR formula (van Regemorter 1962), which is based on the Born and Bethe approximations with an empirical effective Gaunt factor, and the semi-classical impact parameter (IP) method from Seaton (1962). The IP method has been extended to the case of positive ions, where electrons have hyperbolic
trajectories in the semi-classical picture, by Burgess & Summers (1976), but this seems to be little used.

It is well known that the Born approximation overestimates cross-sections at low energies and the IP method gives better results, as was reported for instance by Seaton (1962); see also Bely & van Regemorter (1970) for a discussion. Figure 2 compares results from the two methods for allowed transitions in Mg I. The vR results are larger than those from IP. We also performed tests using either IP or vR in the atomic model, and we found that while we can reproduce the IR Mg I emission lines observed in the Sun when using the IP method, we could not do so when using the vR formula (see Fig. 8). Thus, there is both physical and astrophysical support for preferring the IP method over vR when R-matrix calculations are not available.

Collisional ionisation of low-lying states by electrons was implemented according to the hydrogenic approximation presented in Cox (2000, Sect 3.6.1), which is based on semi-empirical expressions taken from Bely & van Regemorter (1970), which originate from Percival (1966). For Rydberg levels we used the formula from Vrinceanu (2005), where the dependency of the collisional cross-sections on the total electronic orbital angular momentum of the Rydberg atom is taken into account.

R-matrix calculations. We now give the details of the electron collision calculations for the lowest ten states of Mg I using the R-matrix (RM) method. All calculations were performed in L-S coupling. Electronic structure calculations were performed using the code CIVPOL (Hibbert et al. 1977; see also Plummer et al. 2004), which employs the configuration interaction method. CIVPOL is an adaptation of CIV3 (Hibbert 1975), allowing the construction of polarisation pseudo-states. Orbitals for core electrons, 1s, 2s, 2p and 3s, were taken from the ground-state Hartree-Fock calculations of Clementi & Roetti (1974). Valence orbitals, 4s, 5s, 3p, 4p, 3d, 4d and 4f, were optimised with CIVPOL, providing energy levels that agree well with experimental values taken from NIST; see Table 4.

We calculated the excitation cross-sections with the RM method (Burke et al. 1971; Burke & Robb 1976; Berrington et al. 1974, 1978). This method separates the calculation into two cases: when the colliding electron is in the internal region, that is, close to the target atom where interactions are strong, and when the colliding electron is in the external region, far enough from the target atom such that interactions are weak and correlation and exchange effects can be neglected. The R-matrix ensures continuity between wave functions at the

Table 3. Broadening cross-sections, velocity parameters, and line widths for the lines of interest.

| Wavelength [μm] | Transition | \(\sigma^{3\text{h}}\) \(10^{457}\text{ m/s}\) | \(\alpha\) | \(\sigma^{3\text{h}}(10^{457}\text{ m/s})\) \(\text{[au]}\) | \(\gamma\) \(\text{[au]}\) | \(w/N_\text{H}\) \(\text{[cm}^3\text{ rad s}^{-1}]\) |
|-----------------|------------|----------------------|---------|-----------------------------|-------------------|------------------|
| 7               | 5g–6h      | 4950                 | 1.549   | 7.52                        | 2.2               | 13.4             |
| 12.2            | 6g–7h      | 5191                 | 1.738   | 804                         | 48.4              | 13.4             |
| 18.8            | 7g–8h      | 4497                 | 1.764   | 4303                        | 3560              | 14.7             |
| 18.95           | 7h–8i      | 4304                 | 1.776   | 13.7                        | 13.7              | 14.3             |
| 18.96           | 7i–8k      | 3831                 | 1.789   | 12.2                        | 12.2              | 14.3             |

Notes. Columns 3 and 4 give broadening parameters, the cross-section \(\sigma^{3\text{h}}\) at \(v = 10^4\) m/s, and the velocity parameter \(\alpha\) (the ABO theory format). Columns 5 and 6 compare the cross-sections from this work with those of HBvR at \(v = 10^{457}\) m/s. The broadening cross-section can be determined directly from their line width ((\(\gamma H/\gamma H_{\text{H}}\))/2 × 1/(\(\gamma H\)) in their notation). In Cols. 7 and 8 the line widths (half-width at half-maximum) at \(T = 5000\) K are compared.

Table 4. Energies for Mg I states from electronic structure calculations and comparison with experimental values.

| State            | Energy (eV) | Experiment | CIVPOL |
|------------------|-------------|------------|--------|
| 2p^63s^2         | 1S          | –          | –      |
| 3s3p             | 3P          | 2.71       | 2.67   |
| 3s3p             | 1P          | 4.34       | 4.38   |
| 3s4s             | 3S          | 5.11       | 5.10   |
| 3s4s             | 1S          | 5.39       | 5.36   |
| 3s3d             | 1D          | 5.75       | 5.73   |
| 3s4p             | 3P          | 5.93       | 5.88   |
| 3s3d             | 2D          | 5.94       | 6.01   |
| 3s4d             | 1P          | 6.12       | 6.05   |
| 3s5s             | 3S          | 6.43       | 6.43   |

Notes. The energies shown are relative to the ground state of Mg I. The experimental values were taken from NIST (Ralchenko et al. 2010).
boundary between the two regions. To perform the RM calculations for the internal region we used the code RMATRIX I (Berrington et al. 1995), and for the external region a version of the code STGF (Berrington et al. 1987) modified to treat collisions with neutral atoms (Badnell 1999).

After exploring the effects of the input data on the final collisional cross-sections, we adopted a target atom consisting of 25 spectroscopic states (i.e., energy levels), with ten additional pseudo-states states to represent the Rydberg levels and the continuum, giving 35 states in total. Target atoms with more levels were also tested with no significant changes in the resulting cross-sections.

The maximum occupation numbers in each $nl$ orbital used to build the configurations for both $N$ and $N + 1$ atomic structures are shown in Table 5.

The boundary between the internal and external regions was set at 30.4 atomic units. The final calculation included partial waves $L \leq 20$. We made test calculations using partial waves up to $L \leq 30$ to check convergence. $R$-matrix calculations using $L \leq 10$ provided rate coefficients within 20% of those calculated using $L \leq 20$ except for the 3d $^3D$–$^1P^o$ transition, where the rate coefficient increased by a factor of 2. There were no significant differences between the resulting rate coefficients when using $L \leq 30$ and $L \leq 20$ in the temperature range typical of cool stellar atmospheres.

The final collisional cross-sections obtained were compared with experiments and with other calculations where available. Zatsarinny et al. (2009) calculated electron collisional excitation cross-sections for Mg I from the ground state 3s $^1S$ to the excited states 3p $^1P^o$, 3d $^3D$, 4s $^1S$ and 4p $^1P^o$ and compared them with other calculations and with experiments. The cross-sections calculated in this work agree with those within ~20%. Rate coefficients were obtained by integrating the final cross-sections weighted by a Maxwellian velocity distribution and are presented in Table 7.

Figure 3 compares our results with those obtained with the IP and vR formulas. Both approximate formulas typically yield lower values than the RM results. Most past studies of Mg in non-LTE have, however, employed the electron collisional data for transitions between low-lying levels that were presented in Mauas et al. (1988). Mauas et al. studied Mg I non-LTE effects in the Sun using a 13-level model atom, and their compilation of electron collision data, adopting data from different sources (see that paper for details), has been employed extensively since. Thus, it is of interest to compare our results with their compilation, which is done in Fig. 4. Allowed transitions (circles) have results similar to ours, but for forbidden transitions (diamonds) our results are significantly larger. For forbidden transitions Mauas et al. mostly used rates equal to 0.1 times the collisional rate of an allowed transition with a similar energy difference calculated using the vR formula.

For optically forbidden transitions, there are no simple approximate formula similar to vR or IP, and we require some method to estimate the rates of forbidden transitions between highly excited levels. Common approaches to circumvent this problem are to adopt an arbitrary collision strength $\Omega_j$ of the order of unity, or to assume collision strengths similar to nearby optically allowed transitions. We adopted a more nuanced approach. First, from a physical point of view, it makes sense to divide optically forbidden transitions into two groups: those involving electron exchange (i.e. corresponding to intersystem transitions with change of spin state) and those not involving electron exchange. At high energy, transitions not involving electron exchange may be described in the Born approximation, while transitions involving electron exchange are described in the Ochkur approximation. This leads to significantly different behaviour of the cross-sections with energy, at least at high energy (e.g. Bransden & Joachain 2003; Burgess & Tully 1992).
calculations as a function of temperature

Table 6. Average values of $T_{ij}/g_i$ (dimensionless) calculated from RM calculations as a function of temperature $T$.

| $T$ [K] | Non-exchange | Exchange |
|---------|--------------|----------|
| 1000    | 0.086        | 1.020    |
| 2000    | 0.174        | 0.654    |
| 3000    | 0.257        | 0.635    |
| 4000    | 0.336        | 0.642    |
| 5000    | 0.413        | 0.650    |
| 6000    | 0.492        | 0.655    |
| 7000    | 0.568        | 0.658    |
| 8000    | 0.646        | 0.660    |
| 10000   | 0.797        | 0.658    |

Second, we then assume that transitions of the same type (exchange or non-exchange) and same transition energy will have similar rate coefficients. Using Eq. (20) of Burgess & Tully (1992), this implies a constant value of $T_{ij}/g_i$ at a given temperature, where $T_{ij}$ is the thermally averaged collision strength (e.g. Burgess & Tully 1992, Eq. (21)), and $g_i$ is the statistical weight of the initial state $i$.

Indeed, we found using the results of the RM calculations that the behaviour of $T_{ij}/g_i$ for a given transition type was similar, but significantly different between the two groups. Among the calculated RM collisions we have 4 non-exchange forbidden transitions and 40 spin-exchange transitions. We calculated the mean value for $T_{ij}/g_i$ for the two groups from the RM calculations, and the results are presented in Table 6. The scatter of the spin-exchange transitions about the mean value is greatest at low temperatures, becoming smaller at higher temperatures. At 5000 K the values span over three orders of magnitude (0.01 $\leq T_{ij}/g_i \leq 10$) with half of the values within a factor of 2 of the mean. The mean behaviour for exchange transitions with temperature is rather flat (except at the lowest temperature), while for non-exchange transitions the values steadily increase over the temperature range of interest. To estimate rate coefficients for optically forbidden transitions not included in the RM calculations, we interpolate in this table, and then use Eq. (20) of Burgess & Tully (1992), which relates the thermally averaged collision strength to the rate coefficient.

3.3.2. Hydrogen atoms

The rate coefficients, $\langle \sigma v \rangle$, for excitation and de-excitation processes,

$$\text{Mg}(3s n \, l^{2S+1}L) + \text{H}(1s) \rightleftharpoons \text{Mg}(3s n' l'^{2S'+1}L') + \text{H}(1s),$$

(3)

and for the charge transfer processes, ion-pair production and mutual-neutralisation, involving the ionic state,

$$\text{Mg}(3s n \, l^{2S+1}L) + \text{H}(1s) \rightleftharpoons \text{Mg}^+ (3s \, 2S^+) + \text{H}^-, \tag{4}$$

are taken from Barklem et al. (2012), which are based on calculations of Belyaev et al. (2012); see also Guitou et al. (2010, 2011). These ab initio quantum chemical calculations on which the scattering calculations are based were obtained using the code MOLPRO (Werner et al. 2009) and cover the seven lowest lying states of Mg I and the ground state of Mg II.

We wish to emphasize that with regard to hydrogen collisions in general, the accurate quantum (Barklem et al. 2003; Barklem et al. 2010, 2012) as well as the more approximate model calculations (Belyaev 2013; Belyaev et al. 2014), based on a correct physical background, provide non-zero rate coefficients for both optically allowed and optically forbidden transitions, as well as for charge transfer processes, which generally have the highest rate coefficients among all hydrogen collision processes. Thus, the Drawin formula fails to provide reliable estimates for important inelastic processes in collisions with hydrogen. In the case of Mg, the quantum eight-channel calculations Belyaev et al. (2012) provide 28 non-zero cross-sections for the endoergic processes (excitation and ion-pair production), while the Drawin formula gives only five non-zero (overestimated) cross-sections for the same 28 transitions (see Barklem et al. 2012). This is a general feature of the Drawin formula, as discussed in detail by Barklem et al. (2011).

For collisional excitation involving higher lying and Rydberg states of Mg I, we employed the free electron model of Kaulakys (1986, 1991), which considers a binary encounter of the Rydberg electron with a neutral perturber. This approach allows writing the cross-section for the inelastic transition $n \rightarrow n'$ in terms of the elastic differential cross-section for the electron-perturber scattering and the momentum-space wave functions of the initial state (Kaulakys 1991, Eqs. (6)–(8)). For the wave functions $g_\alpha(p)$, where $p$ is the electron momentum, we calculated non-hydrogenic wave functions from quantum-defect theory following Hoang Binh & van Regemorter (1997). For neutral hydrogen perturbers, the relevant elastic differential cross-section $|f_{\alpha}(p, \theta)|^2$ is that for $e^-H$ collisions, where $f_{\alpha}(p, \theta)$ is the scattering amplitude. This total cross-section can be written in terms of the singlet and triplet amplitudes, $f^s$ and $f^t$, which is found by averaging over initial spins and summing over final spins to obtain the well-known expression (e.g. Seaton 1962)

$$|f_{\alpha}(p, \theta)|^2 = \frac{\alpha^2}{4} |f^s(p, \theta)|^2 + \frac{\alpha^2}{4} |f^t(p, \theta)|^2. \tag{5}$$

The elastic singlet and triplet scattering amplitudes for $e^-H$ collisions are known from variational calculations of the phase shifts for $s$- and $p$-wave scattering (Schwartz 1961; Armstead 1968). We make use of the form introduced by HBvR, namely

$$|f^\pm(p, \theta)|^2 = A^\pm + B^\pm \cos(\theta), \tag{6}$$

where $A^\pm$ is the $s$-wave contribution and $B^\pm$ the $s$-$p$ coupling. The coefficients $A^\pm$ and $B^\pm$ are listed as a function of $p$ in Table 3 of HBvR, derived from the published phase shifts mentioned above. From this we calculated the total cross-section for the transition $n \rightarrow n'$, which we denote $\sigma_{n \rightarrow n'}$.

In our model atom, singlet and triplet terms are often separated, and thus we would like to estimate how this total cross-section is distributed among final spin states. We considered a representation where the spin of the Rydberg electron is coupled to the spin of the other electrons in the target atom $S_c$ (assuming this to be a good quantum number) to give the total electronic spin of the Rydberg atom, $S$. Recoupling of angular momenta gives the required spin-changing scattering amplitude in terms of the singlet and triplet scattering amplitudes (i.e. in the representation in which the Rydberg electron spin and the hydrogen atom spin are coupled), and the differential cross-section for change of spin $S \rightarrow S'$, $S \neq S'$, is found to be

$$|f_{\alpha}(p, \theta, S \rightarrow S')|^2 = \frac{(2S' + 1)}{8(2S_c + 1)} |f^s - f^t|^2 \tag{7}$$

and

$$= \frac{(2S' + 1)}{2(2S_c + 1)} \sigma_{\text{exc}}. \tag{8}$$

\footnote{The required Hankel transforms were derived analytically for $l$ ranging from 0 to 13 using Mathematica. We note that the denominators in Eqs. (32) and (33) of Hoang Binh & van Regemorter (1997) contain a misprint where an exponent is incorrectly written as a multiplicative factor - the denominator of Eq. (32) should be $|v^2 + q^2|^{(1+\text{ex})/2}$.}
Fig. 5. Collision matrices for Mg I and Mg II at $\log \tau_{500} = -0.23$ in the solar $(3D_{\odot})_{1D}$ atmospheric model, which corresponds to a temperature of 6000 K. The legend in each case defines a range of $\log C_{ij}$, where $C_{ij}$ is the total collision rate in s$^{-1}$, such that darker squares represent higher rates. Note the 3p$^2$ level of Mg I is not collisionally coupled to other levels; see Sect. 3.3.

where $\sigma_{\text{exch}}$ is the differential cross-section for electron exchange in elastic $e^-+H$ collisions (e.g. Field 1958; Burke & Schey 1962). The derivation essentially follows Dalgarno & Rudge (1965), except that in our case the electron spin is coupled to the core electronic spin and not to the nuclear spin. We note that $S' = S_c \pm 1/2$ and the factor $(2S' + 1)/(2S_c + 1)$ is always in the range 0 to 1 and represents the probability that an electron exchange leads to a change in the total electronic spin of the Rydberg atom. In the limit $p \rightarrow 0$, $-f^\pm \rightarrow a^\pm$, where $a^\pm$ are the scattering lengths, which have values of $a^+ = 5.965$ and $a^- = 1.769$ atomic units (Schwartz 1961). In this limit, for the case of $S_c = 1/2$, via comparison of the expressions for the cross
Table 7. Rate coefficients for de-excitation of Mg I by electron collisions.

| Initial \ Final | 3s 1S | 3p 1P* | 3p 1P* | 4s 1S | 4s 3S | 3d 1D | 4p 3P* | 3d 1D | 4p 1P* |
|-----------------|-------|--------|--------|-------|-------|-------|--------|-------|--------|
|                 |       |        |        |       |       |       |        |       |        |
| 1000 K          |       |        |        |       |       |       |        |       |        |
| 3p 1P           | 1.79E 08 |       |       |        |       |       |        |       |        |
| 3p 1P           | 5.65E 08 | 6.59E 07 |       |        |       |       |        |       |        |
| 4s 1S           | 1.11E 08 | 4.57E 07 | 7.91E 08 |        |       |       |        |       |        |
| 4s 1S           | 7.37E 08 | 1.74E 07 | 2.22E 07 | 1.31E 07 |       |       |        |       |        |
| 3d 1D           | 2.77E 09 | 9.74E 08 | 8.36E 08 | 9.25E 08 | 2.47E 08 |       |        |       |        |
| 4p 1P           | 3.10E 09 | 2.68E 07 | 4.95E 07 | 3.98E 07 | 5.87E 08 | 6.24E 07 |       |       |        |
| 3d 1D           | 8.93E 10 | 6.88E 08 | 1.53E 08 | 9.98E 08 | 1.45E 08 | 1.22E 07 | 4.25E 05 |       |        |
| 4p 1P           | 4.24E 09 | 8.25E 08 | 6.19E 08 | 8.95E 08 | 6.57E 08 | 6.57E 08 | 5.75E 08 | 2.58E 06 | 1.98E 05 |
| 5s 1S           | 1.10E 07 | 5.30E 06 | 1.21E 06 | 5.57E 06 | 2.58E 06 | 1.98E 05 | 4.17E 06 | 2.26E 05 | 6.83E 06 |
| 5000 K          |       |        |        |       |       |       |        |       |        |
| 3p 1P           | 3.18E 08 |       |       |        |       |       |        |       |        |
| 3p 1P           | 6.71E 08 | 3.04E 07 |       |        |       |       |        |       |        |
| 4s 1S           | 1.54E 08 | 2.55E 07 | 8.09E 08 |       |       |       |        |       |        |
| 4s 1S           | 1.45E 08 | 1.18E 07 | 3.67E 07 | 1.42E 07 |       |       |        |       |        |
| 3d 1D           | 6.50E 09 | 8.58E 08 | 1.26E 07 | 7.86E 08 | 3.85E 08 |       |        |       |        |
| 4p 1P           | 3.85E 09 | 1.62E 07 | 4.37E 08 | 3.08E 07 | 3.08E 08 | 2.87E 07 |       |       |        |
| 3d 1D           | 1.38E 09 | 5.31E 08 | 1.98E 08 | 1.42E 07 | 9.67E 09 | 8.60E 08 | 2.24E 05 |       |        |
| 4p 1P           | 8.14E 09 | 6.72E 08 | 9.58E 08 | 3.48E 08 | 3.10E 07 | 1.51E 06 | 1.33E 07 | 1.82E 07 |       |
| 5s 1S           | 2.02E 08 | 4.21E 07 | 1.99E 07 | 4.20E 07 | 2.57E 07 | 1.84E 06 | 6.18E 07 | 1.81E 06 | 1.36E 06 |
| 8000 K          |       |        |        |       |       |       |        |       |        |
| 3p 1P           | 3.19E 08 |       |       |        |       |       |        |       |        |
| 3p 1P           | 6.93E 08 | 2.25E 07 |       |        |       |       |        |       |        |
| 4s 1S           | 1.36E 08 | 1.95E 07 | 6.84E 08 |       |       |       |        |       |        |
| 4s 1S           | 1.51E 08 | 9.31E 08 | 4.21E 07 | 1.24E 07 |       |       |        |       |        |
| 3d 1D           | 7.81E 09 | 7.51E 08 | 1.25E 07 | 5.98E 08 | 4.14E 08 |       |        |       |        |
| 4p 1P           | 3.65E 09 | 1.41E 07 | 3.92E 08 | 2.69E 07 | 2.16E 08 | 2.04E 07 |       |       |        |
| 3d 1D           | 1.38E 09 | 4.53E 08 | 1.84E 08 | 1.85E 07 | 7.41E 09 | 6.64E 08 | 1.66E 05 |       |        |
| 4p 1P           | 1.02E 09 | 6.05E 08 | 1.12E 07 | 2.98E 08 | 4.13E 08 | 1.55E 06 | 1.13E 07 | 1.56E 07 |       |
| 5s 1S           | 1.33E 08 | 2.25E 07 | 1.52E 07 | 1.83E 07 | 1.54E 07 | 1.03E 06 | 3.47E 07 | 8.21E 07 | 1.03E 06 |
| 10000 K         |       |        |        |       |       |       |        |       |        |
| 3p 1P           | 3.04E 08 |       |       |        |       |       |        |       |        |
| 3p 1P           | 7.05E 08 | 1.95E 07 |       |        |       |       |        |       |        |
| 4s 1S           | 1.22E 08 | 1.73E 07 | 6.09E 08 |       |       |       |        |       |        |
| 4s 1S           | 1.49E 08 | 8.22E 08 | 4.44E 07 | 1.13E 07 |       |       |        |       |        |
| 3d 1D           | 8.22E 09 | 6.96E 08 | 1.23E 07 | 5.10E 08 | 4.19E 08 |       |        |       |        |
| 4p 1P           | 3.46E 09 | 1.35E 07 | 3.68E 08 | 2.51E 07 | 1.81E 08 | 1.74E 07 |       |       |        |
| 3d 1D           | 1.33E 09 | 4.21E 08 | 1.72E 08 | 2.10E 07 | 6.50E 09 | 5.75E 08 | 1.44E 05 |       |        |
| 4p 1P           | 1.13E 09 | 5.70E 08 | 1.21E 07 | 2.73E 08 | 4.70E 08 | 1.56E 06 | 1.02E 07 | 1.44E 07 |       |
| 5s 1S           | 1.16E 08 | 1.80E 07 | 1.49E 07 | 1.31E 07 | 1.34E 07 | 8.60E 07 | 2.75E 07 | 5.98E 07 | 9.61E 07 |

Notes. Rate coefficients, \( \langle \sigma v \rangle \), are given in cm³ s⁻¹.

sections we find that

\[
\sigma_{nlS \rightarrow \pi' \pi' \pi'} = 0.706 \sigma_{nl \rightarrow \pi' \pi'} , \\
\sigma_{nlS \rightarrow \pi' \pi' \pi' } = 0.294 \sigma_{nl \rightarrow \pi' \pi'} , \\
\sigma_{nlS \rightarrow \pi' \pi' \pi'} = 0.098 \sigma_{nl \rightarrow \pi' \pi'} , \\
\sigma_{nlS \rightarrow \pi' \pi' \pi'} = 0.902 \sigma_{nl \rightarrow \pi' \pi'} .
\]

3.4. Test model atoms

To understand the uncertainties in the modelling, we constructed various model atoms in which sets of collisional data were changed or removed compared to the final model. For ease

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of discussion, we adopted the following abbreviations for the various collision processes:

| CE | Collisions (de)excitation by electrons |
| C1 | Collisional ionisation by electrons |
| CH | Collisions (de)excitation by hydrogen |
| CH0 | Charge transfer with hydrogen |

ence, our final model adopted for CE data RM data wherever possible, IP data for all remaining allowed transitions, and we used the recipe described in Sect. 3.3.1 and defined by Table 6 for forbidden transitions (hereafter referred to as the “T” recipe). The final model, which we give the label F, also includes data for CI, CH and CH0 as described in Sect. 3.3.

The first additional test model represents a basic model without any of the new data introduced here. This model contains only CE and C1, and no hydrogen collision processes. For CE it employs IP data for all allowed transitions and uses the “T” recipe for all forbidden transitions. Thus, we refer to it, where appropriate, as the basic model, model B.

The models F and B therefore represent two extremes, and it is interesting to investigate models that lie between these cases. In particular, we constructed models B + RM (introduces RM), B + RM + CH (further introduces CH), B + RM + CH0 (introduces RM and CH0), B + H (introduces CH and CH0), which may be used to explore the relative importance of RM, CH, CH0, or hydrogen collisions in total, H, in causing the differences between model B and model F. To isolate the effects of spin changing collisions, we constructed versions of the final model where spin changing collisions (i.e. between singlet and triplet states in Mg I) are removed for collisions with electrons (F−SCE), hydrogen (F−SCH) or both (F−S). Transitions between levels of different spin can only be removed when the spin of each level is defined, but not in merged super levels.

Finally, we constructed slightly modified versions of model F, in which the IP data were replaced with vR data and/or the T recipe was replaced by a typical approximation of setting the collision strength to unity, Ω = 1. These models are labelled F(vR, T), F(vR, Ω = 1), F(IP, Ω = 1), noting that F(IP, T) would be equivalent to the final model F in this notation, and these models can be used to investigate the impact of various approximations for CE among high-lying levels not covered by RM data.

Table 8 gives an overview of these models. By examining all of these models, we can determine for a given model atmosphere which of the newly introduced data or processes has produced the most important changes in each line, and this is reported in the last four columns of Table 1 and is part of our discussions of below, where we discuss the results for the various groups of lines.

### 4. Comparisons with observations

We now test our modelling by comparison with observed spectra. First, we compare our results with optical spectra of standard benchmark late-type stars, including the Sun. Then we compare them with the IR emission lines in the solar spectrum.

#### 4.1. Optical absorption lines

We tested the performance of our non-LTE modelling using high-quality spectra of six stars with reasonably well-known fundamental parameters. This sample of benchmark stars was designed to cover a wide parameter range and includes the Sun, Procyon, Arcturus, and three very metal-poor stars ([Fe/H] = −2.5 ... −2.0); a turn-off star (HD 84937), a sub-giant (HD 142083), and a red giant (HD 122563). For the Sun and Arcturus, spectra from the Kitt Peak observatory with R ≈ 150 000 and S/N ≈ 1000 are available (Hinkle et al. 2000). For the other stars, we retrieved VLT/UVES spectra at R ≈ 47 000 from the POP archive of bright field stars (Bagnulo et al. 2003).

Accurate effective temperatures for Procyon, Arcturus, and HD 122563 were derived from direct measurements of their angular diameters (see adopted parameters in Table 9). Creevey et al. (2015) presented fundamental parameters of HD 140283, although T eff is uncertain, and for HD 84937 no such measurements have been published, so we relied instead on the infra-red flux method (Casagrande et al. 2010). For all stars except for the Sun, HIPPARCOS parallax measurements were used to calculate the surface gravity (van Leeuwen 2007). Reference metallicities were taken from Jofré et al. (2014) for Arcturus and Bergemann et al. (2012) for all other stars. Microturbulence values were also taken from the literature studies and held fixed in the line formation calculations.

We obtained 1D and (3D)IP model atmospheres for each star by interpolating in the MARCS (Gustafsson et al. 2008) and STAGGER (Magic et al. 2013) grids. Minor extrapolation in the (3D)IP grid was sometimes necessary to produce the desired model parameters. The 1D spectrum synthesis code SME (Valenti & Piskunov 1996, and subsequent updates) was used to compute synthetic spectra including blending lines. To synthesise the lines in non-LTE, grids of pre-computed departure coefficients from MULTI (using model F) were read in and interpolated before they were applied to the LTE level populations computed by SME. For all interpolation and line-formation, the continuum optical depth at 5000 Å was used as reference scale, which is consistent with the reference scale used when computing the spatial

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6 These notations are adopted from MULTI.

7 The ‘T’ recipe is a new addition, based on new RM data. However, its importance is limited and we chose to use it in the basic model to limit the number of degrees of freedom in our tests. Its effects are tested via the F(IP, Ω = 1) model described later in this section.

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| Name | Allowed CE | Forbidden CE | CH | CH0 |
|------|------------|--------------|-----|-----|
| B    | F          |              |     |     |
| B + RM | RM + IP    | RM + T       |     |     |
| B + RM + CH | RM + IP    | RM + T       |     |     |
| F vR |                |              |     |     |
| F IP | IP          | T            |     |     |
| F (IP, T) | RM + IP    | RM + T       |     |     |
| F (IP, Ω = 1) | RM + IP    | RM + (Ω = 1) |     |     |
| F (vR, Ω = 1) | RM + vR    | RM + T       |     |     |
| F (vR, Ω = 1) | RM + vR    | RM + (Ω = 1) |     |     |

**Notes.** RM + x means RM when available, otherwise method x.

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8 Defined as bi = ni / ni where ni is the non-LTE population and ni is the LTE population of level i.
and temporal average of the full 3D models. We compared the spectral line profiles calculated by MULTI and SME used in this fashion, and they give practically identical results.

Between eight and eleven Mg lines, selected to be relatively blend-free and situated between 4000–9000 Å were synthesised per star. Whenever possible, the best-fitting abundance was found by simultaneously varying the Mg abundance and the intrinsic line broadening, such as caused by macro-turbulence and rotation. It was often found that neglecting non-LTE effects in the cores of strong lines prevented a match between the LTE synthetic spectrum and the observations without artificially reducing all broadening, even the instrumental, to unphysical values. In such cases, we instead fixed the broadening parameters to the values found in non-LTE and disregarded the line core when optimising the fit. By doing so, the agreement between LTE and non-LTE abundances is typically better than if the same equivalent widths had been enforced, for instance. For lines weaker than 150 mÅ, the LTE and non-LTE line profiles are similar, and a good fit to the full observed line profile can always be found. Sample (3D)_{1D} fits are seen in Fig. 6 for Mg I 8806 Å.

The choice of model atmosphere in 1D or (3D)_{1D} and line-formation method in LTE or non-LTE affects the derived Mg abundances to various degrees and in various directions. Average abundances and line-by-line 1σ dispersions are presented in Table 9. We stress that our main goal is to illustrate the impact of the modelling method on a variety of different lines, not only those that are most suitable for abundance analysis. The

References. Model parameters taken from (60) Bergemann et al. (2012); (60) Jofré et al. (2014).

Table 9. Stellar parameters and average Mg abundances, including line-to-line 1σ dispersions, for the benchmark stars.

| Name          | $T_{\text{eff}}$ | $\log g$ | [Fe/H] | $v_{\text{mic}}$ [km s$^{-1}$] | $\langle \text{Mg}\rangle_{1D}$ | (3D)$_{1D}$ |
|---------------|------------------|----------|--------|-------------------------------|----------------------------------|-------------|
| Sun           | 5777             | 4.44     | 0.00   | 1.09                          | 7.57 (0.08)                      | 7.67 (0.07) |
| Procyon$^a$   | 6545             | 3.99     | −0.03  | 2.00                          | 7.47 (0.04)                      | 7.60 (0.05) |
| Arcturus$^b$  | 4247             | 1.59     | −0.52  | 1.63                          | 7.42 (0.07)                      | 7.52 (0.10) |
| HD14907       | 5777             | 3.67     | −2.40  | 1.23                          | 5.36 (0.10)                      | 5.35 (0.16) |
| HD10283       | 6408             | 4.13     | −2.03  | 1.40                          | 5.73 (0.04)                      | 5.78 (0.05) |
| HD122563$^b$  | 4608             | 1.61     | −2.64  | 1.50                          | 5.29 (0.13)                      | 5.28 (0.08) |

Fig. 6. Observed spectra of the non-LTE sensitive Mg I 8806 Å line (pluses). The best-fitting (3D)$_{1D}$LTE (red dashed line) and non-LTE profile (blue dotted line) are overplotted and the corresponding Mg abundances stated in each panel.
average abundances we present should therefore not be taken too literally. For the Sun, non-LTE effects are very weak, of the order of 0.01 dex, while (3D)_{1D} abundances are higher by 0.05–0.15 dex than the corresponding 1D abundances because of the hotter temperatures (~1%) in the line-formation region. Only two lines overlap with the recent study by Scott et al. (2015), but the average solar abundances nevertheless match well (A(Mg)_{Scott et al.} = 7.59 ± 0.04).

In Arcturus and Procyon the effects work in the same direction as in the Sun, but the sensitivity to non-LTE is higher. The over-population of Mg_{i} in the Sun results from contributions of Mg_{i} in the photosphere with a non-LTE source function in- ffected by the strong cooling of the surface layers that is characteristic of metal-poor radiation-hydrodynamical models (Asplund & García Pérez 2001). Consequently, (3D)_{1D} LTE abundances are significantly lower for these lines, up to ~0.25 dex in HD122563. However, the effects of the non-LTE cooling on line strengths is counteracted by non-LTE effects in the opposite direction. The behaviour is qualitatively similar to that found for resonance lines of Ca_{i} in Lind et al. (2013) and Fe_{i} in Bergemann et al. (2012). As also described in these studies, the steeper temperature gradients of metal-poor (3D)_{1D} models than in 1D models strengthens the overionisation of neutral species, such as Mg_{i}, resulting in positive non-LTE corrections for all lines. Reassuringly, the line-to-line abundance scatter in (3D)_{1D} non-LTE is overall an improvement with respect to 1D LTE.

In Fig. 7 we illustrate how abundances of the benchmark stars compare when limited to the diagnostics available to the GALAH and Gaia-ESO survey for metal-rich and metal-poor stars. Evidently, systematic uncertainties are comparable to or even exceed the targeted precision (0.1 and 0.05 dex for Gaia-ESO and GALAH, respectively). The subgiant and dwarf stars in our sample do not appear to suffer from strong differential effects, and sufficiently precise relative abundances can probably be obtained also with traditional modelling of Mg, although they may not necessarily be accurate in an absolute sense. However, giants show strong differential non-LTE effects (0.1–0.2 dex), which must be corrected for to allow comparisons to less evolved stars.

4.2. Infrared emission lines

As discussed in Sect. 2, the formation of IR Mg_{i} emission lines in the solar spectrum, discovered in the early 80s, took a decade and considerable effort to understand. Two main hypotheses arose: they form in the chromosphere under LTE conditions, or they form in the photosphere with a non-LTE source function increasing outwards. This debate was finally and unambiguously resolved by Carlsson et al. (1992) in favour of a non-LTE line formation scenario of photospheric origin. Their modelling made several reasonable, yet ad hoc, assumptions regarding collision rates. As pointed out by Rutten & Carlsson (1994), the 12 μm lines have non-LTE emissions peaks because of the strong collisional coupling between the pertinent levels. Based on the expectation that hydrogen collisions would provide such high rates (Omont 1977), Carlsson et al. introduced high electron collisional rates for 1-changing collisions among Rydberg states to ensure detailed balancing between levels of equal n. Attempts at modelling since that time, such as Zhao et al. (1998) and Sundqvist et al. (2008), have described hydrogen collisions using the highly debated Drawin formula scaled by a free parameter chosen to best fit observations. Importantly, however, these studies showed that the IR lines in the Sun and in red giants such as Arcturus are rather sensitive to the description of hydrogen collisions, and as these lines probe the highly excited parts of the atom, they are important tests of our ab initio modelling.

Figure 8 compares theoretical spectra for the 7.3, 12.2, and 12.3 μm emission lines with the observed solar spectrum across the solar disk. The synthetic spectral lines are calculated for various model atoms, using the currently accepted solar photospheric Mg abundance (Scott et al. 2015), A(Mg) = 7.59, and employing the (3D)_{1D} solar model atmosphere. We note that our modelling does not include pressure-induced line shifts, and other atomic data for such shifts do not exist. Thus, we have no expectation to be able to model the asymmetries observed in the wings of these lines.

First, we note the well-known and practically obvious result that the lines cannot be reproduced in a photospheric model using LTE line formation. Second, we see that our model F reproduces the observations both in the core and the wings rather well, but certainly not perfectly. However, it certainly reproduces the observations better than other model atoms of interest. As noted in Sect. 3.3.1, on physical grounds the vR formula is expected to overestimate the CE rates because it is based on the Born approximation. The vR formula results in rates higher than those given by the IP method, producing stronger collisional coupling between levels of different n and thus reducing the strength of the emission. The predicted line cores for the model F(vR,T) are in all cases far too weak. In addition, we also see that a model atom with no hydrogen collision processes, model B+RM, produces line profiles that are generally too strong, both in the core and in the wings. This supports the conclusion of earlier studies that additional collisional mechanisms are required. Thus, we see that our model F employing the IP method for CE rates among Rydberg states, and the Kaulakys method for CH among Rydberg states, generally produces the line profiles better than models using the vR method or ignoring CH. The fact that the...
Fig. 8. Solar IR line profiles at various positions on the disk. First row: 7.3 μm line, second row: 12.2 μm line, and final row: 12.3 μm line. The columns are μ = 1.0, 0.5, and 0.2, in that order. The circles are observations taken from Chang et al. (1991) for the 7.3 μm line and from Brault & Noyes (1983) for the two 12 μm lines. The synthetic spectral lines are calculated for various model atoms using \( A(Mg) = 7.6 \) employing the \( \langle 3D \rangle_{1D} \) solar model atmosphere. For the synthetic profiles, a macroturbulent velocity of 3.15 km s\(^{-1}\) and a rotational broadening of \( v \sin i = 1.6 \) km s\(^{-1}\) are used. For model F, the profile without macroturbulence is also shown (grey full line) to demonstrate its effect.

The astrophysical comparison agrees with expectations based on the physical characteristics of the vR and IP methods, giving us confidence that our ab initio modelling of collision rates among Rydberg states due to CE and CH is satisfactory.

We note that the emission peaks are sensitive to macroturbulence (see Fig. 8) and the wings can be better fitted with a change in abundance of order 0.1 dex. As might be reasonably expected, the CH0 processes do not significantly affect these lines because CH0 processes only involve relatively low-lying states, and thus any effect must be indirect.

Sundqvist et al. (2008) studied these lines in a small sample of K giants including Arcturus, in which they showed that the 12.2 μm line was particularly sensitive to the modelling of hydrogen collisions. This is due to the small influence of electron collisions in such atmospheres. Figure 9 compares our results with their observations (kindly provided by Nils Ryde). The LTE
Fig. 9. Profile of the 12.2 μm line observed in Arcturus by Sundqvist et al. (black circles) and compared with our calculations using the (3D)_{1D} atmospheric model and the non-LTE abundance taken from Table 9. The black lines show synthetic profiles using different model atoms (see Table 8), and are all convolved with a macroturbulent velocity of 5.2 km s^{-1} and a rotational broadening of \(\varepsilon\sin i = 2.1\) km s^{-1}. The grey line is the profile using the F model atom but without macroturbulence.

Table 10. Stellar parameters of the atmospheric models used in Sect. 5.

| Name             | \(T_{\text{eff}}\) [K] | \(\log g\) [cm/s^{2}] | [Fe/H] [dex] | \(v_{\text{mic}}\) [km s^{-1}] |
|-------------------|------------------------|------------------------|-------------|-------------------------------|
| Dwarf poor        | 6000                   | 4.5                    | -2.0        | 1.0                           |
| Dwarf rich        | 6000                   | 4.5                    | 0.0         | 1.0                           |
| Giant poor        | 4500                   | 4.0                    | -2.0        | 2.0                           |
| Giant rich        | 4500                   | 4.0                    | 0.0         | 2.0                           |

calculation is again not able to form this line in emission. For our model atom F, the match in the wings is good, while the emission peak is stronger than in the observed spectra; however, we consider the agreement reasonable given, for example, the uncertainties in stellar parameters and that the macroturbulence and abundance have not been fine-tuned. As is the case for the Sun, our model F reproduces the line better or as well as the other models of interest. In particular, neglect of hydrogen collisions, model B+RM, again produces a peak considerably too strong. We note that in this case, the F(vR) model performs marginally better than the F(IP) model; however, the differences are small and we judge that this is insignificant with respect to the other modelling uncertainties mentioned above.

5. Spectral line behaviour

In this section we examine the behaviour of the modelled spectral lines, in particular the influence of different collisional processes on line strengths and the abundances that would be derived from the lines. Tests were performed on a small grid of four 1D model atmospheres from the MARCS grid (Gustafsson et al. 2008) with parameters described in Table 10 where \(v_{\text{mic}}\) is the micro turbulent velocity used in calculating the model atmospheres. The same value is used in the line formation calculations. Additionally, we performed a number of test calculations for the solar case, using the MARCS solar model. We also performed calculations using the photospheric reference model with a chromosphere from Maltby et al. (1986), hereafter MACKKL, and the 1D-averaged 3D model for the Sun, (3D)_{1D}, from Magic et al. (2013). Furthermore, to investigate possible effects of the chromosphere, we also made a hybrid model using MACKKL at \(\tau_{\text{500nm}} > -3\), while replacing the upper layers with those from the (3D)_{1D} model. Where comparisons are made with solar spectra, we used a Gaussian macro-turbulence with \(v_{\text{mic}} = 2.0\) km s^{-1}.

As we are most interested in effects on the derived abundance of Mg, we studied the influence of the various collision processes on the line formation of Mg lines through the non-LTE corrections, defined by

\[
\Delta a_{\text{Mg}}^{\text{NLTE-LTE}} = A_{\text{Mg}}^{\text{NLTE}} - A_{\text{Mg}}^{\text{LTE}}. \tag{13}
\]

The abundance corrections discussed in this section are based on matching equivalent widths. We note that they therefore cannot be compared to the changes in abundances seen in the last section, where profile fitting was used.

For the lines to be discussed here, various non-LTE mechanisms interact to varying degrees to produce the final line profiles. Generally, overionization leads to an underpopulation of the low-lying levels of Mg I, and a number of Mg lines lead to photon losses, resulting in departures from LTE also among intermediate levels (e.g. Carlsson et al. 1992). These radiative mechanisms may be counteracted or even propagated by collisional processes. Charge exchange processes (CH0) drive the populations of specific intermediate levels, in particular those around 4s1S, toward detailed balance with the ground state of Mg II, which is generally close to LTE populations. CH couples low-lying and intermediate levels propagating departures from low-lying to higher levels.

5.1. Mg I lines

3829, 3832, and 3838 Å (3D → 3p): The UV triplet is one of the strongest features of Mg I and is often used for abundance determinations in metal-poor stars. In solar metallicity stars these lines are significantly blended, and also sensitive to chromospheric effects. The abundance corrections and their sensitivity to collisional processes are presented in Fig. 10. Model B leads to very large corrections for these lines. However, in model F abundance corrections are general quite small. In dwarfs the corrections are practically non-existent, but if collisions with H were not included, we would find corrections of \(\sim 0.2\) dex in the metal-poor case. Thus, in dwarfs CH and CH0 play an important role in the abundance correction calculation. In giants, corrections are also small, \(\lesssim 0.05\) dex for abundance values of interest, and neglecting CH leads to much larger corrections, of about 0.2 dex in the metal-poor case. This means that in giants, CH strongly contributes to the abundance correction and CH0 has a negligible effect. If hydrogen collisions are included, the use of more approximate models for CE, such as F(vR, \(\Omega = 1\)) or F(IP, \(\Omega = 1\)) does not change the results significantly. Spin change transitions have only a weak effect in the two dwarf stellar models, the correction changing by \(\sim 0.05\) dex in the metal-poor case for the F–S model. In giants, the F–S model changes the corrections by \(\sim 0.07\) dex with the relative effect of SCE being strongest at solar metallicity and the effect of CH0 strongest at low metallicity.
Fig. 10. Abundance corrections for the 3838 Å, 5528 Å, 4571 Å, 5184 Å, and 8718 Å lines in four different atmospheric models and using modified model atoms as described in the text. The model parameters are given at the bottom left of each figure with $T_{\text{eff}}$/$\log g$/$[\text{Fe}/H]$. The shaded region indicates the likely abundance of Mg for a normal star of given metallicity. The black circles show the calculated abundances; some lines have fewer points because the calculations with those parameters did not converge. Note that the model atoms shown in the 4571 Å line case differ from the others.

4167, 4703, 5528, and 8806 Å (nd 3D–3p 1P°): for a given stellar model, all four of these lines behave similarly, but the effects become stronger at longer wavelengths, where the lines form at shallower depths with larger population departures. Figure 10 shows the abundance corrections for the 5528 Å line. Model B leads to large corrections in dwarfs, of about 0.2 dex at the abundances of most interest. The corrections with model F are, however, small. The results for 4167, 4703, and 5528 were most sensitive to the introduction of hydrogen collisions generally (both CH and CH0), while for the 8806 line CH was particularly important. This is because the populations of the levels involved are similarly affected by the introduction of CH0; the upper level of the 8806 Å transition, 3d 1D, is low-lying and both levels are thermalized by CH0 to similar degrees in the line formation region, thus leaving $S_{\nu}$ unaltered.

4571 Å (3p 3P°–3s 1S): the intercombination line, despite its weak oscillator strength, is observable across a wide range of stellar parameters because it involves the ground state of Mg i. The abundance corrections are shown in Fig. 10. In dwarfs, at abundances of typical interest, this line is formed close to LTE.
conditions as the collisional coupling of the two levels in different spin systems dominates the weak radiative coupling. In giant models where collisional coupling is significantly decreased, this line shows significant departures from LTE, of the order of 0.2 dex at solar metallicity and almost 0.4 dex in the metal-poor giant model. In metal-rich giants, CH0 is much weaker than either CH and RM. This line is expected to be sensitive to spin change transitions. CE rates between the upper and lower levels are 108 times higher than the corresponding CH rates, making this line most sensitive to spin change CE rates even in very metal-poor stars.

5167, 5172, and 5184 Å (4s3S–3p3P)\(^i\): the abundance corrections for one of the Mg I \(^i\) triplet lines are shown in Fig. 10. Model B gives large corrections for these lines, but model F gives corrections that are very small for dwarfs, and quite small for giants, lower than 0.05 dex. As with the 8806 Å line, the departure coefficients of the upper and lower levels are similarly affected by CH0, and thus CH0 has little effect on the line strength. The main change is due to the introduction of CH. CH decreases the population of the upper level in the line formation region and increases the populations of the lower levels, bringing them closer to relative LTE.

8736 Å (7d\(^3\)F–3d\(^3\)D) and 8710, 8713, and 8718 Å (7d\(^3\)D–4p\(^3\)P)\(^i\): these lines are \(\tau = 7 \leftrightarrow \tau' = 3\) transitions involving triplet states, that is, between low-lying and high-lying levels. The abundance corrections for one of the lines are shown in Fig. 10. These lines show examples of behaviour where the addition of collision processes can drive away from LTE, rather than towards it. As these levels lie approximately 3–4 eV from the continuum, they are subject to overionisation from ultraviolet radiation. In the solar case, Carlsson et al. (1992) have shown that the triplet 3d and 4p levels are among those involved in transitions that become optically thin in the photosphere, which in turn leads to photon losses and thus photon suction. Now, together with these radiatively driven non-LTE processes, the CH and CH0 processes involving these states are among the strongest.

The complex interplay of these processes leads to the situation seen in Fig. 10, where rather than the more usual convergence from model B towards the F model, the introduction of additional collisional processes has less predictable effects. For example, in the metal-poor dwarf case, with model B, rather large corrections are seen. Introduction of the RM data via model B+RM leads to slightly increased corrections. Then, contrary to the naive expectation that additional collisions drive towards LTE, the introduction of CH, model B+RM+CH actually increases the abundance corrections. The left panel of Fig. 11 shows departure coefficients of the levels involved in the formation of the 8718 Å line in a metal-poor giant atmosphere, where we see that the introduction of CH leads to populations of the 4p level at around \(\log(\tau_{500 \text{ nm}}) = -2\) to \(-3\) closer to the LTE population, but the population at around \(-1\), where the line forms, is in fact farther from LTE. This occurs because deep in the atmosphere, CH processes lead to increased coupling with the lower-lying levels, especially 4s and 3p levels, which are subject to strong photoionisation and thus violent overionisation, leading to underpopulation. However, we see in this model atmosphere that the dominant change is due to the introduction of CH0, which increases the coupling between the 4s, 3d and 4p levels with the continuum and thus brings the population closer to relative equilibrium with Mg II and thus towards LTE populations, particularly in the line-forming region. For the metal-poor giant model, with departure coefficients shown in the right panel of Fig. 11, model B shows an overpopulation of the 4p level in the line-forming region \(\log(\tau_{500 \text{ nm}}) = -1\) caused by photon losses. The introduction of CH processes again pulls the populations towards those of the lower-lying levels, which are underpopulated between \(\log(\tau_{500 \text{ nm}}) = -1\) and 0.5. This causes an underpopulation at large depths, but brings the population to near LTE levels in the line-forming region. Introduction of CH0 processes brings the population deep in the atmosphere back towards LTE, which in turn reintroduces a slight overpopulation in the line-forming region.

5.2. Mg II lines

Since Mg II is the dominant ion of Mg in late-type stars, non-LTE effects are expected to be weaker in Mg II than in Mg I lines. In the four test model atmospheres studied here, the abundance corrections all had magnitudes lower than 0.1 dex for the lines we analysed (see Table 1).
6. Conclusions

We have presented a new model atom for Mg suitable for non-LTE line formation studies in late-type stars. It includes data for electronic collision rates calculated in this paper by the $R$-matrix method, and we have further used these results to formulate a method to estimate transition rates for forbidden transitions not covered by the $R$-matrix calculations, and not possible to be estimated by the vR and IP methods. Recent data for hydrogen collisions, including charge transfer processes, calculated by some of us, were also applied. Some new data for collisional broadening were calculated as well. The new model atom, used together with 1D model atmospheres in the context of standard non-LTE modelling, compares well with observations. The modelled spectra agree well with observed spectra from benchmark stars, showing much better agreement with line profile shapes than LTE modelling. The line-to-line scatter in the derived abundances shows some improvements compared to LTE (where the cores of strong lines must often be ignored), particularly when coupled with averaged 3D models. The observed Mg emission features at 7 and 12 μm in the spectra of the Sun and Arcturus, which are sensitive to the collision data, are reasonably well reproduced.

The spectral line behaviour and uncertainties were explored by extensive experiments in which sets of collisional data were changed or removed on a small grid of theoretical model atmospheres. Charge transfer in collisions with H is generally important as a thermalising mechanism in dwarfs, less so in giants. Excitation due to collisions with H was found to be quite important in both giants and dwarfs. The $R$-matrix calculations for electron collisions also lead to significant differences compared to when approximate formulas are employed. The modelling predicts non-LTE abundance corrections $\Delta(A(Mg)_{\text{NLTE}})$ in dwarfs, both solar metallicity and metal-poor, to be very small (of about 0.01 dex), even smaller than found in previous studies. In giants, corrections vary greatly between lines, but can be as large as 0.4 dex. Our results emphasise the need for accurate data for electron and hydrogen collisions for precise non-LTE predictions of stellar spectra, but demonstrate that such data can be calculated and that ab initio non-LTE modelling without resort to free parameters is possible.

Only two other elements, Li and Na, have collisional data for non-LTE modelling of comparable quality to that which has been used in this work. RM calculations have been performed for excitation of low-lying levels of Li and Na by electron collisions by Osorio et al. (2011) and Gao et al. (2010), respectively. Before calculating the RM data, the data of Park (1971) were used in non-LTE modelling, and these data typically agree to within 80% of the RM data, significantly better than the vR or IP methods. Thus, the new RM data for electron collisions did not have a strong effect on non-LTE modelling, since the existing data were relatively good; see Osorio et al. (2011) for Li, Lind et al. (2011) for Na. Data for hydrogen collision processes, including charge transfer, have been calculated for Li by Barklem et al. (2003), and for Na by Barklem et al. (2010), and used in non-LTE studies by Barklem et al. and Lind et al. (2009) in the case of Li and Lind et al. (2011) for Na. These studies found for both elements that CH processes have practically no effect on the non-LTE modelling, while CH0, in particular that involving the first excited $S$-state, had a significant effect because it brought this level into closer detailed balance with the continuum. These are, however, alkalis with only one spin system. In contrast, Mg has two spin systems, leading to the added feature of intersystem transitions. Furthermore, many more lines of Mg are observable in stellar spectra from IR to UV. Our modelling has shown that the situation for slightly more complex atoms such as Mg is far more complicated than for alkalis, with the new data for $CE$, CH, and CH0 all affecting at least some lines.

We plan to use the current model atom to produce grids of departure coefficients and abundance corrections in 1D and (3D)$_{1D}$ theoretical model atmospheres, specifically the MARCS (Gustafsson et al. 2008) and STAGGER (Magie et al. 2013) grids. Grids of abundance corrections can be directly applied to studies performed in LTE. However, this depends on the use of equivalent widths, which is only strictly useful for weak lines. Grids of departure coefficients allow non-LTE corrections to be made at the spectrum synthesis level, and thus is far more robust when using stronger lines. We have demonstrated that non-LTE calculations are necessary for Galactic archaeology studies with large spectroscopic surveys of the Milky-Way. In particular, Mg abundances of giant stars suffer from strong differential non-LTE effects with respect to dwarfs and subgiants, comparable to or exceeding the targeted precision. Using grids of departure coefficients computed with our model atom, on-going and future surveys can properly model these effects.

Finally, we note that a comparison with centre-to-limb variations of optical lines in the Sun would provide a strong complement to the tests of the modelling performed here. Such an observational campaign is underway with the Swedish Solar Telescope and will be the subject of future work. Ideally, this should be done in the context of full 3D non-LTE modelling.

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