Three-dimensional hydrodynamical CO$^5$BOLD model atmospheres of red giant stars

III. Line formation in the atmospheres of giants located close to the base of the red giant branch

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ABSTRACT

Aims. We utilize state-of-the-art three-dimensional (3D) hydrodynamical and classical 1D stellar model atmospheres to study the influence of convection on the formation properties of various atomic and molecular spectral lines in the atmospheres of four red giant stars, located close to the base of the red giant branch, RGB ($T_{\text{eff}} \approx 5000$ K, log $g = 2.5$), and characterized by four different metallicities, [M/H] = 0.0, −1.0, −2.0, −3.0.

Methods. The role of convection in the spectral line formation is assessed with the aid of abundance corrections, i.e., the differences in abundances predicted for a given equivalent width of a particular spectral line with the 3D and 1D model atmospheres. The 3D hydrodynamical and classical 1D model atmospheres used in this study were calculated with the CO$^5$BOLD and 1D LHD codes, respectively.

Identical atmospheric parameters, chemical composition, equation of state, and opacities were used with both codes, therefore allowing a strictly differential analysis of the line formation properties in the 3D and 1D models.

Results. We find that for lines of certain neutral atoms, such as Mg I, Ti I, Fe I, and Ni I, the abundance corrections strongly depend both on the metallicity of a given model atmosphere and the line excitation potential, $\chi$. While abundance corrections for all lines of both neutral and ionized elements tend to be small at solar metallicity ($\pm 0.1$ dex), for lines of neutral elements with low ionization potential and low-to-intermediate $\chi$ they quickly increase with decreasing metallicity, reaching in their extremes −0.6 to −0.8 dex. In all such cases the large abundance corrections are due to horizontal temperature fluctuations in the 3D hydrodynamical models. Lines of neutral elements with higher ionization potentials ($E_{\text{ion}} \geq 10$ eV) generally behave very similarly to lines of ionized elements characterized by low ionization potentials ($E_{\text{ion}} \leq 6$ eV). In the latter case, the abundance corrections are small (generally, $\pm 0.1$ dex) and are caused by approximately equal contributions from the horizontal temperature fluctuations and differences between the temperature profiles in the 3D and 1D model atmospheres. Abundance corrections of molecular lines are very sensitive to the metallicity of the underlying model atmosphere and may be larger (in absolute value) than $\sim 0.5$ dex at [M/H] = −3.0 ($\sim 1.5$ dex in the case of CO). At fixed metallicity and excitation potential, the abundance corrections show little variation within the wavelength range studied here, 400–1600 nm. We also find that an approximate treatment of scattering in the 3D model calculations (i.e., ignoring the scattering opacity in the outer, optically thin, atmosphere) leads to abundance corrections that are altered by less than $\sim 0.1$ dex, both for atomic and molecular (CO) lines, with respect to the model where scattering is treated as true absorption throughout the entire atmosphere, with the largest differences for the resonance and low-excitation lines.

Key words. stars: atmospheres – stars: late-type – stars: abundances – convection – hydrodynamics

1. Introduction

All low and intermediate mass stars evolve through the red giant stage. Owing to their large numbers, red giants are important tracers of intermediate-age and old stellar populations, and their high luminosities make them accessible for observations both in and beyond the Milky Way. This makes them particularly suitable for use in spectroscopic studies that aim to understand the chemical evolution of their host populations. On the other hand, stars on the red giant branch (RGB) experience mixing episodes during which material with the chemical composition altered via nuclear reactions is brought to the stellar surface. Elemental abundances in the atmospheres of red giant stars may therefore provide a wealth of important information about
the nucleosynthesis and mixing processes in stars and about the chemical evolution histories of their host populations.

To obtain information from stellar spectra about the abundances of the chemical species of interest one has to rely on stellar model atmospheres. Inevitably, the confidence with which one can derive chemical composition of stars is determined by the accuracy and physical realism of the underlying model atmospheres. Classical 1D model atmospheres are most widely used for such tasks today. Stellar atmospheres in these models are treated as static and 1D, so that resulting model structures are functions of the radial coordinate alone. Convection in such models is described by using the mixing-length theory (MLT; see, e.g., Böhm-Vitense 1958; Canuto & Mazzitelli 1991), along with a number of free parameters, such as the mixing-length parameter, \( \alpha_{\text{MLT}} \), and microturbulence velocity. Thanks to their static 1D nature and various simplifications involved, such models are fast to compute, which allowed large model grids to be produced using a variety of 1D model atmosphere codes, such as ATLAS (Castelli & Kurucz 2003), MARCS (Gustafsson et al. 2008), and PHOENIX (Brott & Hauschildt 2005), for various combinations of stellar parameter space. Despite numerous simplifications, 1D model atmospheres have aided in gaining enormous amounts of information about the processes taking place in stellar atmospheres and deeper interiors.

Since stellar atmospheres are neither static nor 1D, a large increase in computational power is required to include more realistic 3D time-dependent physics into stellar atmosphere modeling. Mostly because of such computational complexities the advance of 3D hydrodynamical model atmospheres and their applications in stellar abundance work has been relatively slow. Despite this, there has been a notable increase in the applications of the 3D hydrodynamical model atmospheres to study stars and stellar populations during the recent years (see, e.g., Asplund et al. 2000; Caffau et al. 2008; González Hernández et al. 2009; Behara et al. 2010). This is partly related to the fact that a number of studies have shown that differences in the elemental abundances inferred from the same spectral line strength with the classical 1D and 3D hydrodynamical model atmospheres may be significant, especially at lower metallicities where they may reach \( \sim 1.0 \) dex (see, e.g., Collet et al. 2007; Dobrovolskis et al. 2010; Ivanovauskas et al. 2010). Obviously, such differences in the obtained abundances may lead to important implications, such as when reconstructing most likely scenarios of chemical evolution of stellar populations in the Galaxy and beyond.

To take a closer look at the role of convection in the spectral line formation in the atmospheres of red giant stars, we started a systematic study of the internal structures and observable properties of red giant stars by using 3D hydrodynamical CO5BOLD model atmospheres for this purpose. In the first two papers in this series (Ludwig & Kučinskas 2012; Kučinskas et al. 2013, Papers I and II hereafter), we investigated the role of convection on the atmospheric structures and spectral line formation in a red giant star located close to the RGB tip (\( T_{\text{eff}} \approx 3600 \) K, \( \log g = 1.0 \), \( [\text{M/H}] = 0.0 \)). In this study we extend our previous work and focus on the properties of spectral line formation in somewhat warmer red giants at several different metallicities (\( [\text{M/H}] = 0.0, -1.0, -2.0, -3.0 \)) located close to the RGB base (\( T_{\text{eff}} \approx 5000 \) K, \( \log g = 2.5 \)).

The paper is structured as follows. In Sect. 2 we describe the 3D hydrodynamical and classical 1D model atmospheres used in this study, and outline the details of spectral line synthesis calculations. In this section we also introduce the concept of abundance corrections, which is used throughout the paper to assess the differences in the abundances predicted for the same line strength with the 3D and 1D model atmospheres. The abundance corrections obtained for various neutral and ionized atoms and for molecules are discussed in Sect. 3, together with the role of scattering and the importance of the mixing-length parameter, \( \alpha_{\text{MLT}} \), used with the 1D model atmospheres. Finally, summary and conclusions are provided in Sect. 4.

## 2. Stellar atmosphere models and spectral line synthesis calculations

In this study we utilized the CO5BOLD and LHD model atmosphere codes to calculate, respectively, the 3D hydrodynamical and classical 1D atmosphere models of red giants located close to the RGB base. Both CO5BOLD and LHD simulations shared identical atmospheric parameters, chemical composition, opacities, and equation of state, in order to limit the discrepancies in the model predictions to differences in the underlying model physics. Such differential comparison of the model predictions allows us to assess the importance of 3D hydrodynamical effects on the spectral line formation, and enables us to determine the differences between the abundances of different chemical elements derived with the 3D and 1D model atmospheres. In what follows below we briefly describe the details of model atmosphere and spectral line synthesis calculations.

### 2.1. 3D hydrodynamical CO5BOLD stellar model atmospheres

The 3D hydrodynamical models used in this study are part of the CIFIST model atmosphere grid which will eventually cover stars on the main sequence, subgiant, and red giant branches (Ludwig et al. 2009). CO5BOLD solves the coupled nonlinear equations of compressible hydrodynamics in an external gravity field, together with radiative transfer equation (Freytag et al. 2012). The model atmospheres were computed using the “box-in-a-star” setup, i.e., the part of the stellar atmosphere modeled was small compared to the size of a star itself. The simulations were performed on a Cartesian grid of \( 160 \times 160 \times 200 \) grid points in \( x, y, z \) directions, respectively. The model box had open upper and lower boundaries (matter was allowed to enter and leave the simulation box freely), and periodic boundaries in the horizontal direction (matter leaving the box on one side was allowed to enter it again from the opposite side).

We used monochromatic opacities from the MARCS stellar atmosphere package (Gustafsson et al. 2008) which we grouped into a smaller number of opacity bins using the opacity binning technique (Nordlund 1982; Ludwig 1992; Ludwig et al. 1994; Vöglér 2004; Vöglér et al. 2004), with five opacity bins for the [M/H] = 0.0 model and six bins for the [M/H] = −1.0, −2.0, and −3.0 models. Solar-scaled elemental abundances used were those from Asplund et al. (2005). It is important to stress that we also applied a constant enhancement in the alpha-element abundances of \( \alpha/\text{Fe} = +0.4 \) for the models at metallicities [M/H] \( \leq -1.0 \). All model simulations were performed under the assumption of local thermodynamic equilibrium, LTE. Scattering was treated as true absorption and magnetic fields were neglected.

Parameters of the individual 3D model atmospheres used in this work are provided in Table 1. One may notice that each model is characterized by a slightly different average effective temperature, \( \langle T_{\text{eff}} \rangle \). This is because effective temperature is not the input parameter for calculating 3D model atmosphere with the CO5BOLD code. Instead, one sets the value of entropy
Table 1. Parameters of the CO$^3$BOLD red giant models used in this work.

| $(T_{\text{eff}})_i$ | $\log g_i$ | [M/H] | Grid dimension, Mm | Resolution, grid points |
|---------------------|-----------|-------|-------------------|------------------------|
| 4970                | 2.5       | 0     | $573 \times 573 \times 243$ | $160 \times 160 \times 200$ |
| 4990                | 2.5       | -1    | $573 \times 573 \times 245$ | $160 \times 160 \times 200$ |
| 5020                | 2.5       | -2    | $584 \times 584 \times 245$ | $160 \times 160 \times 200$ |
| 5020                | 2.5       | -3    | $573 \times 573 \times 245$ | $160 \times 160 \times 200$ |

2.2. 1D LHD model atmospheres

1D models were calculated using the LHD code (see, e.g., Caffau et al. 2008). We recall that both 3D and 1D models used in this work shared identical atmospheric parameters (Table 1), chemical composition, equation of state, and opacities (see Sect. 2.1 for details). Convection in the LHD models was treated according to the formalism of the mixing-length theory by Mihalas (1978). The mixing-length parameter used throughout the LHD simulations was $\alpha_{\text{M.L.}} = 1.0$. In principle, the use of 1D models calculated with different mixing-length parameters results in different abundance corrections. However, in the case of the giants studied here it is only relevant for high-excitation atomic lines (see Sect. 3.5). Temperature stratifications of the 3D and 1D models at $[\text{M/H}] = 0.0$ and $-3.0$ are shown in Figs. 1 and 2, respectively.

2.3. Spectral line synthesis

In order to investigate the influence of convection on the spectral line formation we utilized fictitious spectral lines. For these lines, their central wavelengths, excitation potentials, $\chi$, and oscillator strengths, $\log gf$, were freely chosen to cover the range of values of existing spectral lines. Such approach allows us to study the connections between the atomic line parameters and line formation properties in the presence of realistically modeled convective core and convective envelope. We have already applied successfully in a number of studies (e.g., Steffen & Holweger 2003; Collet et al. 2007; Dobrovolskas et al. 2010; Ivanauskas et al. 2010; Paper II).

In this study, equivalent widths of synthetic spectral lines were fixed to $W \leq 0.5$ pm. Because these lines are very weak, they should be confined to the linear part of the curve of growth and thus their strengths should be independent of the choice of microturbulence velocity, $\xi_{\text{mic}}$, used in the spectral line synthesis with the (3D) and 1D model atmospheres. In such situation, the differences in the line strengths predicted by the 3D and 1D model atmospheres can be attributed to purely 3D hydrodynamical effects, allowing investigation of their importance in the spectral line formation.

Spectral line synthesis calculations for all chemical elements studied in this work were done using the LINFOR3D$^1$ code, which solves the radiative transfer equation under the assumption of LTE. In order to speed up spectral line synthesis calculations, we used only every third point of the original 3D model structure along the $x$ and $y$ spatial directions. We verified that such simplification has left the resulting elemental abundances essentially unchanged (i.e., with respect to the calculations performed using the full model box). The differences were always below 0.003 dex in the case of $[\text{M/H}] = -3$ metallicity model and Fe I lines. In our simulations the radiative transfer was solved for 3 vertical and 4 azimuthal directions. Line synthesis calculations covered the Rosseland optical depth range from $\log \tau_{\text{Ross}} = -6$ to 2, in steps of $\Delta \log \tau_{\text{Ross}} = 0.08$. Line profiles were computed with a typical wavelength resolution of 130–150 points per profile. In case of the 3D model atmospheres, spectral line profiles of every chemical element studied here were calculated for each of the twenty 3D model snapshots at each of the four metallicities. The obtained individual spectral line profiles were then co-added to obtain the final 3D line profile at each metallicity. Similarly, in case of the average 3D models line profiles were calculated for every horizontally averaged 3D model structure from the 20-snapshot ensemble; the final (3D) line profile was then obtained by co-adding all twenty individual line profiles.

1. http://www.aip.de/~mst/linfor3D_main.html
All 1D line profiles were also calculated using the Linfor3D code. A microturbulence velocity of $\xi_{\text{mic}} = 1.0 \, \text{km} \, \text{s}^{-1}$ was used in all (3D) and 1D spectral synthesis calculations, although we note again that the choice of $\xi_{\text{mic}}$ does not change the line strength of very weak fictitious lines studied here.

One may envision that, alternatively, all 3D model snapshots could be averaged to produce a single (3D) model for the synthesis of (3D) spectral lines, i.e., instead of using individual averaged 3D models for each 3D snapshot as we did in this study. Our test calculations showed, however, that in case of Fe I and Fe II lines with $\chi = 0–6 \, \text{eV}$ differences in the line profiles computed using these two different approaches would lead to differences in the elemental abundances of less than 0.03 dex (in case of Fe I lines they were always smaller than 0.01 dex). Therefore, for the purposes of present study the choice in the procedure used to compute the (3D) line profiles is of minor importance.

2.4. 3D–1D abundance corrections

The role of 3D hydrodynamical effects in spectral line formation was studied by utilizing 3D–1D abundance corrections. The 3D–1D abundance correction, $\Delta_{3D-1D}$, is the difference in the abundance, $A(X_i)$, of a chemical element $X_i$ obtained at a given equivalent width from the 3D and 1D curves of growth, $\Delta_{3D-1D} \equiv A(X_i)_{3D} - A(X_i)_{1D}$ (see, e.g., Caffau et al. 2011).

The 3D–1D abundance correction is sensitive to the effects related to both the horizontal temperature fluctuations and differences in temperature stratifications of the 3D and 1D model atmospheres. The size of these effects can be measured separately, by utilizing information content of the (3D) models. Since the (3D) model does not contain information about the horizontal fluctuations of thermodynamic quantities and velocity fields, the $\Delta_{3D-1D} \equiv A(X_i)_{3D} - A(X_i)_{1D}$ correction can be used to measure the effect of horizontal temperature fluctuations. Similarly, the $\Delta_{3D-1D} \equiv A(X_i)_{3D} - A(X_i)_{1D}$ correction provides information about the role of differences between the temperature profiles of the average (3D) and 1D models. Obviously, the final abundance correction is a sum of the two constituents, $\Delta_{3D-1D} \equiv \Delta_{3D-1D}^\text{3D} + \Delta_{3D-1D}^\text{1D}$.

3D–1D abundance corrections were calculated for a number of neutral and singly ionized elements: Li I, C I, O I, Na I, Mg I, Al I, Si I, S I I, K I, Ca I, Ca II, Ti I, Ti II, Fe I, Fe II, Ni I, Zn I, Zr I, Zr II, S I I, and Eu II. A sample of very weak fictitious lines was selected for sample different neutral ions: proton capture (Na, Al, K), triple-$\alpha$ (carbon), a-chain captures (O, Mg, Si, Ca, Ti), nuclear statistical equilibrium (Ti, Fe, Ni, Zn), slow neutron capture (Zn, Zr, Ba), rapid neutron capture (Ba, Eu). Lithium is a special element in this context, in a sense that it was produced during the Big-Bang nucleosynthesis and by several other processes afterwards, including cosmic ray spallation in the interstellar medium and several stellar processes. We also obtained abundance corrections for several molecules: CH, CO, C$_2$, NH, CN, and OH which are important tracers of CNO elements.

The 3D–1D abundance corrections for neutral and ionized atoms were calculated at three wavelengths, $\lambda = 400$, 850 and 1600 nm. The two shorter wavelengths were chosen to bracket the typical blue and red limits of modern optical spectrographs, such as UVES/GIRAFFE@VLT, HIRES@Keck, HDS@SUBARU, while the longer approximately coincides with the $H$-band of near-infrared spectrographs, such as CRIRES@VLT, NIRSPEC@Keck. The two longest wavelengths also coincide with the maximum and minimum absorption of the H$^+$ ion at ~850 and ~1640 nm, respectively, while the bluest marks the spectral region where H$^+$ absorption becomes progressively smaller and opacities from various metals become increasingly more important. In case of the molecules, we used the wavelengths of real molecular lines/bands in the blue part of the spectrum (see Paper II for detailed description of the wavelength selection). In case when real molecular bands were not available at these short wavelengths, abundance corrections were obtained at $\lambda = 400$ nm instead. The abundance corrections were computed at four excitation potentials, $\chi = 0$, 2, 4, and 6 eV, for both neutral and ionized atoms, while for the molecules they were obtained at $\chi = 0$, 1, 2, and 3 eV.

Finally, we note that some species studied here have either only a few observable spectral lines or their lines may be simply too weak to be detected (as it is the case with, e.g., Li I, C I, O I). We should stress that such elements were included in the present analysis because we aimed to understand the general properties of spectral line formation in the presence of convection, especially the trends of abundance corrections in a wider range of atomic line parameters. Obviously, in all such cases the exact values of abundance corrections in a wide atomic parameter range may only be of academic interest.

3. Results and discussion

3.1. Abundance corrections for lines of neutral atoms

The 3D–1D abundance corrections for neutral atoms are plotted in Figs. 3–5 for $\lambda = 400$, 850, and 1600 nm, respectively. Each figure shows three abundance corrections, $\Delta_{3D-1D}^\text{3D}$, $\Delta_{3D-1D}^\text{1D}$, and $\Delta_{3D-1D}$, plotted versus metallicity at four different line excitation potentials, $\chi = 0$, 2, 4, and 6 eV.

For some elements, one may notice a strong dependence of the abundance corrections on metallicity: corrections are small at $[\text{M/H}] = 0.0$ and $-1.0$ but they grow quickly with decreasing metallicity and for certain elements may reach up to $-0.8$ dex at $[\text{M/H}] = -3.0$ (Figs. 3–5). Abundance corrections are largest at the lowest excitation potentials, $\chi = 0$, 2 eV, but they quickly decrease with increasing $\chi$: corrections then become both small (less than $\pm 0.1$ dex) and essentially independent of metallicity. Such behavior is defined by the atomic properties of chemical elements and the location of line formation regions associated with particular spectral lines. At all metallicities, lines with lower excitation potentials form in the outer atmospheric layers, but their formation regions shift deeper into the atmosphere with increasing $\chi$ (Figs. 1–2). At solar metallicity, differences between the temperature profiles of the (3D) and 1D model atmospheres are small and change very little throughout the entire model atmosphere. Similarly, horizontal temperature fluctuations, as measured by their rms value ($\Delta T_{\text{rms}} = \sqrt{\langle (T - T_0)^2 \rangle}_{x,y,z}$), here the angle brackets denote temporal and horizontal averaging on surfaces of equal optical depth, and $T_0 = (T/T_{\text{eff}})$, is the depth-dependent average temperature) are not large either (i.e., compared with their extent at lower metallicities) and change little with optical depth (Fig. 1). Therefore, at solar metallicity all three abundance corrections, $\Delta_{3D-1D}^\text{3D}$, $\Delta_{3D-1D}^\text{1D}$, and $\Delta_{3D-1D}$, are small and nearly independent of the line excitation potential. On the other hand, differences in the temperature profiles of the (3D) and 1D models are larger in the outer atmosphere of low metallicity models (Fig. 2). Horizontal temperature fluctuations are also largest in the outer atmosphere, besides, they increase rapidly with decreasing metallicity. Consequently, at $[\text{M/H}] < -1.0$ the abundance corrections for most elements are largest for low-excitation lines, i.e., those that form farthest in the atmosphere. For such lines, $\Delta_{3D-1D}$ correction is significantly
larger than $\Delta_{3D-1D}$, especially at lowest metallicities. Also, the two abundance corrections are nearly always of opposite sign, thus the absolute value of the total correction, $|\Delta_{3D-1D}|$, is smaller than the sum $|\Delta_{3D-3D}| + |\Delta_{3D-1D}|$.

One may notice however, that the size of $\Delta_{3D-1D}$ corrections at a given low metallicity may be very different for different elements, ranging roughly from $-0.8$ dex to $+0.1$ dex at [M/H] = $-3.0$ (Figs. 3–5). Such differences are caused by the interplay of ionization and excitation. Elements with small ionization potential (such as Li I, Na I, K I) are nearly completely ionized throughout the entire atmosphere, at all metallicities. Therefore, neutral atoms of such elements are in their minority ionization stage. In all such cases the line opacity, $\kappa_L$, can be roughly approximated as $\kappa_L \sim 10^\theta (E_{\text{ion}} - \chi)^{-1}$, where $\theta = 5040/T$ and $E_{\text{ion}}$ is the ionization energy of a given element (see Paper II, Appendix A, Eq. (A.5)). In such cases, the line opacity becomes a very sensitive function of the temperature at low $\chi$. Consequently, for lines with low $\chi$ large temperature fluctuations in the outer atmospheric layers at low metallicities cause large variations in the line strength, which translate into large abundance corrections $\Delta_{3D-3D}$, and thus $\Delta_{3D-1D}$. On the other hand, neutral elements with high $E_{\text{ion}}$, such as C I, O I, are in their majority ionization stage. In such case, the excitation potential dominates over the ionization energy and thus the high-excitation lines become most sensitive to the temperature variations. In fact, the dependence of all abundance corrections for C I and O I on both [M/H] and $\chi$ is very similar to those of ionized elements.

Abundance corrections in the infrared wavelength range are very similar to those obtained either at $\lambda = 400$ or $850$ nm and for certain elements may reach up to $\Delta_{3D-1D} = -0.7$ dex. This is in contrast to the results obtained by us for significantly cooler red giant at $T_{\text{eff}} = 3600$ K, $\log g = 1.0$, and [M/H] = 0.0 for which the abundance corrections at 1600 nm were significantly smaller than those in the optical wavelength range (Paper II). Obviously, the latter does not hold for the red giants studied here since their abundance corrections are large at all wavelengths (see Appendix B for a detailed discussion). These results therefore suggest that the use of 3D hydrodynamical models may in fact be necessary for the abundance diagnostics with lines of neutral atoms in red giants at low metallicities.

3.2. Abundance corrections for lines of ionized atoms

The 3D–1D abundance corrections for ionized atoms are shown in Figs. 6–8, at $\lambda = 400, 850$, and 1600 nm, respectively. As in the case with neutral atoms, we provide three abundance corrections, $\Delta_{3D-3D}$, $\Delta_{3D-1D}$, and $\Delta_{3D-1D}$, plotted versus metallicity at four different line excitation potentials, $\chi = 0, 2, 4, 6$ eV.

At all [M/H] and $\chi$ studied here, the abundance corrections for lines of ionized atoms are confined to the range of $\pm 0.1$ dex and show little sensitivity to changes in both metallicity and excitation potential (i.e., if compared to trends seen with lines of neutral atoms). Lines of ionized atoms form significantly deeper in the atmosphere where both the horizontal temperature fluctuations (which determine the size of $\Delta_{3D-3D}$ correction) and the differences between temperature profiles of the (3D) and 1D models (which influence the size of $\Delta_{3D-1D}$ correction) are smallest at all metallicities and change little with [M/H]. This leads to small abundance corrections that are insensitive to changes in [M/H] and $\chi$. On the other hand, elements with lower ionization energies ($E_{\text{ion}} < 6$ eV) are nearly...
completely ionized throughout the entire atmosphere of red giants studied here. For lines of such ionized elements it is the excitation potential that determines the line opacity, \( \kappa \), and thus the strengths of high-excitation lines are most sensitive to temperature fluctuations (see Paper II, Appendix B). Since temperature fluctuations are in fact smallest at the depths where such lines form, and because this holds at all metallicities, this too leads to \( \Delta_{3D-1D} \) corrections that are small and show little variation with either metallicity or excitation potential (though, as expected, they increase slightly with \( \chi \)).

It is worthwhile noting that \( \Delta_{3D-1D} \) and \( \Delta_{3D-1D} \) corrections are often of opposite sign, especially at the lowest metallicities, thus their sum leads to somewhat smaller total abundance corrections, \( \Delta_{3D-1D} \). Since \( \kappa \) is a highly nonlinear function of temperature, horizontal temperature fluctuations produce larger line opacities leading to stronger lines and thus, negative \( \Delta_{3D-1D} \) corrections. On the other hand, since the temperature of the (3D) model is generally lower than that of the 1D model in the line forming regions, lines of ionized species are weaker in (3D) than in 1D. This leads to positive \( \Delta_{3D-1D} \) corrections. As in the case of neutral atoms, abundance corrections at \( \lambda = 1600 \) nm are comparable to those obtained at 400 and 850 nm and may reach to \(-0.1 \) dex at \([M/H] = -3.0\).

Qualitatively, the dependence of abundance corrections on metallicity and excitation potential seen in Figs. 3–7 for neutral and ionized atoms is very similar to the trends found by Collet et al. (2007), who computed abundance corrections for red giants with atmospheric parameters nearly identical to those used in our study. One obvious difference between the results obtained in the two studies is that our abundance corrections are somewhat smaller. It is possible, however, that this discrepancy may be traced back to differences between the underlying 3D model atmospheres, i.e., the STAGGER code used by Collet et al. (2007) and the \textsc{co2bold} code utilized in our study. Indeed, the two codes use different opacities and opacity binning techniques, different equations of state, and so forth. One should also note a major difference in the 1D model atmospheres used as reference. While LHD models use the same opacities, microphysics, and numerical schemes used by \textsc{co2bold}, the \textsc{marcs1d} model atmosphere used as reference by Collet et al. (2007) uses opacities and EOS different from those used in their 3D model.

### 3.3. Abundance corrections for molecular lines

The 3D–1D abundance corrections for molecular lines are shown in Figs. 9, 10, and 11, at the bluest wavelength, 850 nm, and 1600 nm, respectively. We provide three abundance corrections, \( \Delta_{3D-1D} \), \( \Delta_{3D-1D} \), and \( \Delta_{3D-1D} \), plotted versus metallicity at four different excitation potentials, \( \chi = 0, 1, 2, \) and \( 3 \) eV.

As in the case of abundance corrections for neutral atoms, corrections for the molecules are strongly dependent on metallicity and are largest at the lowest metallicities. While for the majority of molecules the total abundance correction, \( \Delta_{3D-1D} \), generally does not exceed \(-1.0 \) dex at \([M/H] = -3.0\), in the case of CO it reaches \(-1.5 \) dex. The large total abundance corrections are mostly due to the large \( \Delta_{3D-1D} \) corrections, which, as in the case of neutral and ionized atoms, indicates the importance of horizontal temperature fluctuations in the formation of molecular lines. The contribution due to differences in the average 3D and 1D temperature profiles, however, is also non-negligible, somewhat in contrast to the much less important \( \Delta_{3D-1D} \) abundance corrections in the case of neutral and ionized atoms.

The trends in the abundance corrections for molecules seen in Figs. 9–11 are determined by a subtle interplay of a number of different factors (see Appendix C for a more
Fig. 7. Same as in Fig. 3 but for lines of ionized atoms, plotted versus metallicity and excitation potential at λ = 850 nm. Abundance corrections at λ = 400 nm and λ = 1600 nm (Figs. 6 and 8, respectively) are available in the online version only.

Fig. 10. Same as in Fig. 3 but for molecular lines plotted versus metallicity and excitation potential at λ = 850 nm. Abundance corrections at λ = 400 nm and λ = 1600 nm (Figs. 9 and 11, respectively) are available in the online version only.

extended discussion). To briefly summarize, as in the case of the cool red giant studied in Paper II CO is the most abundant molecule in the atmospheric layers just above the optical surface (Fig. C.1). Since most of the carbon is locked into CO in the cooler regions (T ≤ 4000 K), only minor amounts of C are available to form other carbon-bearing molecules. In fact, this makes their number densities a highly nonlinear function of the local temperature. Since the line opacity is directly proportional to the number density of a given molecule, large horizontal temperature fluctuations (especially in the lowest metallicity models) should potentially lead to substantial abundance corrections.
experiences the broadest range of fluctuations in the 3D model ([M/H] = −3.0). This is due to the fact that CO has the highest dissociation energy, \(D_0 = 11.1\) eV. Moreover, CO forms rather high in the atmosphere (Fig. C.2) where the largest horizontal temperature fluctuations are encountered. Consequently, the line opacity of CO experiences the largest variations in the line forming region, such that the average line opacity, \(\langle \kappa_T(T) \rangle\) is significantly larger than the line opacity corresponding to the average temperature \(\kappa_T(T)\), leading to the largest (most negative) abundance corrections.

The other molecules, CN (\(D_0 = 7.8\) eV), NH (\(D_0 = 3.5\) eV) and OH (\(D_0 = 4.4\) eV) occupy an intermediate position in the range of abundance corrections. NH and OH have similar molecular properties and line formation regions, and hence their corrections are similar, too. However, it is surprising that the corrections for NH and OH fall in the same range as those for CN, which has a much higher dissociation energy. Again, the explanation is related to the change of sign of the slope \(\partial \log \kappa_T/\partial \theta\) is more positive for \(\Theta = \) than for \(\Theta = \), in accordance with the different values of \(D_0\). But in the low-temperature regions \(\theta > 1.3\), the slope is more negative for \(\Theta = \) than for \(\Theta = \), because \(X(C)\) depends on \(X^2(C)\), while \(X(CH)\) depends only linearly on \(X(C)\). This compensatory effect apparently leads to very similar effective amplification factors (see Appendix B) \(\Delta_{\text{ff}} = \langle \kappa_T(T) \rangle/\kappa_T(T)\) for both molecules, and hence to similar abundance corrections.

These results qualitatively agree with those obtained by Collet et al. (2007), in a sense that the magnitude of abundance corrections for the molecules increases significantly with decreasing metallicity. However, quantitatively the results are different: abundance corrections at the lowest metallicity probed in our study ([M/H] = −3.0) are on average by 0.5–0.6 dex smaller compared with the corrections derived by Collet et al. (2007) using a model with similar atmospheric parameters. For example, for NH molecular line at \(\lambda = 336\) nm and \(\log \chi = 0\) excitation potential we obtain \(\Delta_{\text{1D–3D}} \approx 0.5\) dex while Collet et al. (2007) gets \(\Delta_{\text{1D–1D}} \approx 1.0\) dex. Apparently, these discrepancies are due to differences in the temperature structure of the 3D models used in the two studies. Indeed, in the case of the Stein & Nordlund (1998) models used by Collet et al. (2007) the difference between the average (3D) and 1D temperature stratifications may reach \(-1000\) K at \(\log \tau_{\text{Ross}} \approx -4\) for the [M/H] = −3.0 model, whereas in our case these differences do not exceed \(-300\) K at \(\log \tau_{\text{Ross}} < -1\) (Fig. 2). We would like to stress once more that Collet et al. (2007) compare their 3D results with those obtained using 1D MARS models; clearly, such approach is less self-consistent since differences in, e.g., equation of state, opacities used with the two model atmospheres may affect the resulting 3D–1D abundance corrections. The reason for the different temperature structures is presently not well understood but some possible reasons are discussed below.
The CO\textsuperscript{5}BOLD models treat scattering as true absorption which may be seen as rather crude approximation. Indeed, as it was discussed recently by Collet et al. (2011), the treatment of scattering may have a significant impact on the thermal structure of 3D model atmospheres. In their study, coherent isotropic scattering was implemented in the model atmosphere code and the obtained model structures were compared with those where continuum scattering was treated either as true absorption or scattering opacity neglected in the optically thin regions. These tests have shown that in the former case the resulting temperature profiles were significantly warmer with respect to those calculated with coherent isotropic scattering. It has been thus argued by Collet et al. (2011) that the different thermal structures obtained with the STAGGER and CO\textsuperscript{5}BOLD codes may be due to the different treatment of scattering. Interestingly, the authors also found that in case when scattering opacity was neglected in the optically thin regions the resulting temperature profile was very similar to that obtained when scattering was properly included into the radiative transfer calculations.

The role of scattering in the CO\textsuperscript{5}BOLD model atmospheres has been recently studied by Ludwig & Steffen (2012) who compared thermal structures of the standard CO\textsuperscript{5}BOLD models (i.e., those calculated with continuum scattering treated as true absorption) and the CO\textsuperscript{5}BOLD models computed with continuum scattering opacity left out in the optically thin regions. Surprisingly, temperature profiles in the two models were different by only ~120 K at $\tau_{Ross} = -4.0$, in contrast to ~600 K obtained by Collet et al. (2011). While the exact cause of this difference is still unclear, Ludwig & Steffen (2012) have suggested that they may be due to different procedures used to compute binned opacities used with the two types of models.Obviously, it is important to understand the consequences that the differences in the treatment of scattering may have on the temperature stratification in the model atmosphere which, in turn, may influence the spectral line formation. We therefore calculated a number of fictitious lines of several chemical elements (Li\textsc{i}, O\textsc{i}, Na\textsc{i}, Fe\textsc{i}, Fe\textsc{ii}, Ni\textsc{i}, and Ba\textsc{ii}) by utilizing the CO\textsuperscript{5}BOLD model in which scattering opacity was neglected in the optically thin regions (taken from Ludwig & Steffen 2012). According to the reasoning provided in Collet et al. (2011), the thermal structure of such models should be very similar to that in the models calculated with an exact treatment of scattering. Therefore, the comparison of line formation properties in these and standard models (i.e., those in which scattering is treated as true absorption) may allow to assess the importance of indirect effects of scattering on the spectral line formation via its influence on the temperature profiles. Since scattering becomes increasingly more important at low [M/H] where both line and continuum opacities are significantly reduced, the lowest metallicity CO\textsuperscript{5}BOLD model was used for these tests ($T_{\text{eff}} = 5020$ K, log $g = 2.5$, and [M/H] = ~3.0). Spectral line synthesis was performed using 20 fully relaxed 3D snapshots of this test model, using the procedure identical to that utilized with the standard CO\textsuperscript{5}BOLD models (see Sect. 2.3). The abundance corrections obtained with this and the standard model are shown in Fig. 13. The obtained results suggest that differences in the treatment of scattering within the CO\textsuperscript{5}BOLD setup have only minor influence on the spectral line strengths. The abundance corrections obtained for the standard CO\textsuperscript{5}BOLD model and the one in which scattering opacity was neglected in the optically thin regions differ by less than 0.1 dex, both for neutral atoms and ions (see Fig. 13). For models with scattering opacity neglected, slightly lower temperature in the outer atmospheric layers (Fig. 12) leads to somewhat larger abundance corrections for the low-excitation spectral lines of neutral atoms (~0.1 dex).

Fig. 13. Abundance corrections obtained for selected elements and for the CO molecule with the model atmospheres computed using different treatments of scattering (plotted versus the excitation potential $\lambda$, for lines at $\lambda = 400$ nm). For each element, two cases are shown: (a) with scattering treated as true absorption (solid lines), and (b) with scattering opacity neglected in the optically thin regions (dashed lines). Atmospheric parameters of the model atmospheres are $T_{\text{eff}} = 5020$, log $g = 2.5$, and [M/H] = ~3.0. Three types of abundance corrections are shown: $\Delta_{3D-(3D)}$ (left column), $\Delta_{3D-1D}$ (middle column), and $\Delta_{1D-1D}$ (right column).
for the abundance diagnostics, different recipes in the treatment of scattering may lead to systematic abundance differences of up to 0.1 dex at [M/H] = −3.0. On the other hand, strengths of high-excitation spectral lines, as well as lines of ionized elements, seem to be little affected by the choice of scattering prescription.

3.5. Influence of the mixing-length parameter $\alpha_{\text{MLT}}$ on the abundance corrections

According to the Schwarzschild criterion for the onset of convection, LHD models predict that at Solar metallicity convective flux should be zero at around and above the optical depth unity (Fig. 14, top panel). The situation is slightly different at [M/H] = −3.0 where, because of the lower opacity, convection in the LHD models reaches into layers above the optical surface, with slightly different extension for different choices of the mixing-length parameter, $\alpha_{\text{MLT}}$ (Fig. 14, bottom panel). The majority of spectral lines used in the abundance analysis have $\chi \leq 4$ eV and typically form in the atmospheric layers above log $\tau_{\text{Ross}}$ = 0.0. Such lines should therefore be insensitive to the choice of $\alpha_{\text{MLT}}$, especially at solar metallicity. However, certain exceptions may occur in case of lines characterized by very high excitation potential that form at log $\tau_{\text{Ross}}$ ~ 0.0 or slightly below.

To check the influence of the choice of $\alpha_{\text{MLT}}$ used with the classical 1D models on the abundance corrections, we therefore made several test calculations using LHD model atmospheres computed with $\alpha_{\text{MLT}} = 1.0, 1.5, 2.0$. Abundance corrections were computed for several weak ($W < 0.5$ pm) fictitious lines of Fe I ($\chi = 0$ and 6 eV) and Fe II ($\chi = 6$ eV), at [M/H] = 0.0 and −3.0. The results obtained show that in the case of Fe I lines the dependence on $\alpha_{\text{MLT}}$ at solar metallicity is indeed negligible, with the difference of abundance corrections for $\alpha_{\text{MLT}} = 1.0$ and 2.0 of less than 0.01 dex (−0.04 dex for Fe II $\chi = 6$ eV line, Fig. 15). These differences are somewhat larger at [M/H] = −3.0 but in any case they are below −0.04 and −0.07 dex for Fe I lines with $\chi = 0$ and 6 eV, respectively. The differences are very similar for Fe II lines, too.

The variations in the abundance corrections with $\alpha_{\text{MLT}}$ occur because the temperature profiles of the 1D models at different $\alpha_{\text{MLT}}$ are slightly different at the optical depths where these spectral lines form. For example, temperature in the LHD model with $\alpha_{\text{MLT}} = 1.0$ is slightly higher than in the model with $\alpha_{\text{MLT}} = 2.0$ at the optical depths where the Fe II lines form. This leads to stronger Fe II lines in the model with $\alpha_{\text{MLT}} = 1.0$. Since the lines computed with the 3D models are generally stronger than those obtained in 1D, stronger 1D lines at $\alpha_{\text{MLT}} = 1.0$ lead to slightly less negative abundance corrections with respect to those at $\alpha_{\text{MLT}} = 2.0$.

These test results indicate that the choice of the mixing-length parameter used with the comparison 1D model atmospheres may be important in case of weak, higher-excitation spectral lines, which has also been discussed for the Sun by Caffau et al. (2009). One may expect that in case of stronger lines this dependence may become less pronounced, because such lines tend to form over a wider range of optical depth and typically extend into outer atmospheric layers which are insensitive to the choice of $\alpha_{\text{MLT}}$ used in the 1D models. The results obtained here nevertheless indicate that this issue should be properly taken into account when computing abundance corrections for the higher-excitation spectral lines, especially at lower metallicities.

Since higher excitation lines form deeper in the atmosphere where differences in the thermal profiles are smaller, the influence of differences in the treatment of scattering becomes negligible for such lines, with changes in the abundance corrections of less than 0.01 dex at $\chi > 4$ eV. Lines of ionized elements form deep in the atmosphere too, thus, irrespective of their excitation potential, differences in the treatment of scattering do not affect their line strengths.

The situation may be slightly different in case of molecular lines. The formation of such lines extends into the outer atmosphere where the differences in the treatment of scattering may have an impact on the resulting line strengths. Indeed, in case of CO the differences in the abundance corrections components $\Delta_{3D-1D}$ and $\Delta_{1D-1D}$ calculated using the two treatments of scattering may reach ~0.2 dex and ~0.3 dex, respectively (Fig. 13). However, due to their opposite sign the full 3D–1D abundance correction $\Delta_{3D-1D}$ becomes smaller, ~0.1 dex. Nevertheless, such differences may still be important in stellar abundance work.

Our results therefore suggest that the treatment of scattering may be important in case of the low-excitation lines with $\chi \leq 2$ eV and molecular lines. For elements such as sodium, where frequently only resonance or low-excitation lines are available.
We have studied the influence of convection on the spectral line formation in the model atmospheres of red giant stars located in the lower part of the RGB ($T_{\text{eff}} \approx 5000$ K, log $g = 2.5$), at four different metallicities, [M/H] = 0.0, −1.0, −2.0, −3.0. As in our previous studies (e.g. Dobrovolskas et al. 2010; Ivanauskas et al. 2010, Paper II), the influence of convection was studied by focusing on the 3D–1D abundance corrections, i.e., the differences in abundances predicted for the same line equivalent width by the 3D hydrodynamical CO$^5$BOLD and classical 1D LHD model atmospheres. Abundance corrections were computed for a set of fictitious spectral lines of various astrophysically important neutral and ionized elements (Li i, C i, O i, Na i, Mg i, Al i, Si i, Si ii, K i, Ca i, Ca ii, Ti i, Ti ii, Fe i, Fe ii, Ni i, Zn i, Zr i, Zr ii, Ba ii, and Eu ii), at three different wavelengths ($\lambda = 400$, 850, and 1600 nm) and four line excitation potentials ($\chi = 0, 2, 4, 6$ eV). Only weak lines ($W < 0.5$ pm) were used in the analysis in order to avoid the influence of the microturbulence velocity used with the average (3D) and 1D model atmospheres.

Abundance corrections for the low-excitation lines of neutral atoms show a significant dependence on both metallicity and line excitation potential, especially at low metallicities where differences in the abundances obtained with the 3D and 1D model atmospheres, $\Delta_{3D-1D}$, may reach up to $-0.8$ dex. The corrections are largest for elements with the lowest ionization potentials (generally, $E_{\text{ion}} < 6$ eV), i.e., those that are mostly ionized throughout the entire model atmosphere. The line opacity of the neutral atoms, $\kappa_{\lambda}$, in this case is proportional to $\kappa_{\lambda} \sim 10^{-2} (E_{\text{ion}}/\chi - 3)$ which makes the low-excitation lines most temperature sensitive. Since the formation regions of low-excitation lines extend well into the outer atmosphere where temperature fluctuations are largest, this, in combination with the strong temperature sensitivity of the low excitation lines, leads to the largest abundance corrections. Note that differences between the average temperature profile of the 3D model and that of the 1D model are essentially depth-independent and typically do not exceed a few hundred K, irrespective of the metallicity. The abundance corrections arising due to these differences, $\Delta_{3D-1D}$, are modest, typically $\leq +0.1$ dex, whereas corrections due to horizontal temperature fluctuations, $\Delta_{3D-3D}$, may be significantly larger and reach $-0.8$ dex, which may lead to large total abundance corrections, $\Delta_{1D-1D}$. The corrections decrease quickly with increasing excitation potential and for lines with $\chi > 2$ eV they are confined to $\pm 0.1$ dex within the entire metallicity range. This is because lines with higher $\chi$ are less sensitive to temperature and, besides, they form deeper in the atmosphere where temperature fluctuations are significantly smaller. On the other hand, there is very little variation in the abundance corrections with both metallicity and excitation potential for the lines of neutral atoms of elements with high ionization potentials, i.e., those that are predominantly neutral throughout the entire atmosphere. In this case, it is the high-excitation lines that are most temperature sensitive, but since they form deep in the atmosphere where temperature fluctuations are smaller the resulting abundance corrections are also small.

In case of lines of ionized atoms, the abundance corrections are small at all metallicities and excitation potentials ($\leq +0.1$ dex) and show little variation with either [M/H] or $\chi$. For elements that are predominantly ionized it is the high-excitation lines that are most temperature sensitive. Such lines form deep in the atmosphere where the horizontal temperature fluctuations are small, this leads to small $\Delta_{3D-3D}$ abundance corrections. Since their $\Delta_{3D-1D}$ corrections are never large, the total corrections are therefore significantly smaller than those for lines of, e.g., neutral atoms with low ionization potentials.

Abundance corrections of molecular lines generally depend strongly on the metallicity of the underlying model atmosphere and may reach $\sim -1.0$ dex at [M/H] = −3.0 ($\sim -1.5$ dex in case of CO). Since molecular lines are frequently used to study abundances of astrophysically important elements (such as CNO, for example), the results obtained here indicate that the usage of 3D hydrodynamical model atmospheres in such studies may be essential.

The obtained abundance corrections show little variation with wavelength (both for atoms and molecules), at least in the range of 400–1600 nm. This is in contrast to what was obtained in case of cooler red giant located close to the RGB tip, for which the corrections at 1600 nm were significantly smaller than in the optical wavelength range (Paper II). This may indicate that for certain combinations of atmospheric parameters the effects of convection on the spectral line formation can be equally important over a wide wavelength range.
Rather surprisingly, we find that differences in the treatment of scattering in the radiative transfer calculations seem to have a rather small impact on the resulting thermal structures of the red giant atmospheres studied here, at least within the CO\(^7\)BOLD model setup. The differences in the temperature profiles obtained when scattering is neglected in the optically thin regions and when scattering is treated as true absorption are always confined to \( < 120 \) K and only occur at the optical depths \( \tau_{\text{Ross}} \leq 2.0 \). For certain low-excitation lines this may lead to modest changes in the abundance corrections that are largest for lines of neutral elements with low ionization energies and for the CO molecule, reaching up to \( \sim 0.1 \) dex. For lines of other neutral elements, as well as of ionized species, the changes are significantly smaller (<0.02 dex).

We also find that for the weak, lower-excitation lines (\( \chi < 6 \) eV) there is little dependence of the abundance corrections on the mixing-length parameter, \( \alpha_{\text{MLT}} \) used with the comparison 1D LHD models, both at solar and subsolar metallicities. Differences in the case of high-excitation lines (such as Fe\(^{II}\)) at \( \chi = 10 \) eV are, however, non-negligible and may reach to 0.1 dex. While such sensitivity to \( \alpha_{\text{MLT}} \) may in fact be smaller in case of stronger lines (which form over a wider range of optical depths and reach into the outer atmosphere which is unaffected by the choice of \( \alpha_{\text{MLT}} \) used), these findings nevertheless give an indication that proper care has to be taken to account for the sensitivity on \( \alpha_{\text{MLT}} \) when calculating abundance corrections for high-excitation lines at low metallicities.

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Fig. 3. Same as in Fig. 4 but at $\lambda = 400$ nm.

Fig. 5. Same as in Fig. 4 but at $\lambda = 1600$ nm.
Fig. 6. Same as in Fig. 7 but at $\lambda = 400$ nm.

Fig. 8. Same as in Fig. 7 but at $\lambda = 1600$ nm.
Fig. 9. Same as in Fig. 10 but at $\lambda = 400$ nm.

Fig. 11. Same as in Fig. 10 but at $\lambda = 1600$ nm.
Appendix A: Excitation-like temperature in the (3D) model?

The referee raised the question why an average temperature based on its forth moment is suitable for constructing the (3D) model for comparison. Arguably, for representing spectral properties an average representative of the excitation properties related to the line formation might be the most suitable choice. In this section we want to emphasize that the choice of a suitable average is far from obvious, and our actual choice falls into the range of plausible choices – besides the argument brought already forward that an average based on the forth moment conserves the total flux well.

One may construct an excitation-like temperature $T_e$ in each layer representing the mean population number of a particular energy level produced by the temperature distribution in this layer via

$$T_e^2 \equiv \langle T^b e^{-\chi} \rangle,$$  \hspace{1cm} (A.1)

where $\chi$ is the lower level energy, $k$ is Boltzmann’s constant, and $b$ is an arbitrary exponent. The angular brackets $\langle \rangle$ denote the spatial average. Relation (A.1) is motivated from the Saha-Boltzmann equations governing the excitation and ionization in LTE. In this case, $b \approx 3/2$ or $b = 5/2$ (depending on whether the electron density or electron pressure stays constant during temperature changes), and $\chi$ might have to be identified with the difference between ionization energy and actual level energy in case one deals with a minority species. Factors related to partition functions are ignored. We want to derive an analytical estimate of $T_e$. To this end, we consider the temperature fluctuation around the mean $\overline{T}$ as small and derive the relation to lowest non-trivial order. Introducing for abbreviation the parameter $a \equiv T/\overline{T}$ we obtain for small $x$ ($x = \Delta T/\overline{T}$) the expansion:

$$(1 + x)^b e^{-\chi x} = e^{-a} \left[ 1 + (a + b)x + \frac{1}{2} (a^2 + 2a(b - 1) + b(b - 1)) x^2 \right] + O(x^3).$$

Introducing $\Delta T \equiv T - \overline{T}$ and $\Delta T_e \equiv T_e - \overline{T}$ we obtain as second order

$$\langle T^b e^{-\chi} \rangle \approx \overline{T}^b e^{-\chi \overline{T}} \left[ 1 + \frac{1}{2} \left\{ \left( \frac{\chi}{kt} \right)^2 + \frac{2}{kT} \left( b - 1 \right) \right\} \frac{\Delta T_e}{\overline{T}} \right].$$

$\sigma^2_e \equiv \langle \Delta T^2 \rangle$ is the variance of the temperature distribution. There is no linear term in $\Delta T$ since $\langle \Delta T \rangle = 0$. Similarly, we obtain as first order

$$T_e^2 \equiv \overline{T}^2 e^{-\chi \overline{T}} \left[ 1 + \left( \frac{\chi}{kT} + b \right) \frac{\Delta T_e}{\overline{T}} \right].$$

Combining the expansions (A.3) and (A.4) results in

$$\Delta T_e \approx \left( \frac{\chi}{kT} + (b - 1) - \frac{1}{1 + b / \overline{T}} \right) \sigma^2_e \overline{T}.$$  \hspace{1cm} (A.5)

As an example, for $b = 0$ (corresponding to the level population of a majority species) and $\frac{\chi}{kT} > 2$ the excitation temperature is always greater than the (arithmetic) mean temperature in a given layer. Taking the Sun as a typical example where $kT \approx 0.5$ eV in the line forming region, this means that for lines with $\chi > 1$ eV it follows that $T_e > \overline{T}$. However, the difference between effective excitation temperature and mean temperature is generally not large since the difference depends on the square of the temperature fluctuations.

Average temperatures based on higher moments of the temperature also lead to changes with respect to the mean temperature $\overline{T}$. One may ask which moment is necessary to produce a temperature equal to the effective excitation temperature. With the auxiliary formula

$$(1 + x)^n \approx 1 + nx + \frac{1}{2} n(n - 1)x^2 + O[x^3]$$

we obtain

$$\langle T^n \rangle = \overline{T} \left( \left( 1 + \frac{\Delta T}{\overline{T}} \right)^n \right) \approx \overline{T} \left( 1 + \frac{1}{2} n(n - 1) \sigma^2_e \overline{T} \right),$$

and

$$T^n_e \approx \overline{T} \left( 1 + n \Delta T_e \overline{T} \right) \approx \overline{T} \left( 1 + n \Delta T_e \overline{T} \right).$$

Since we demand $\langle T^n \rangle = T^n_e$ we arrive at

$$\Delta T_e \approx \frac{1}{2} (n - 1) \sigma^2_e \overline{T}.$$  \hspace{1cm} (A.9)

Combining the above result with Eq. (A.5) we obtain for the temperature moment $n$

$$n = \frac{\chi}{kT} + b - \frac{1}{1 + b / \overline{T}} \geq \frac{\chi}{kT} + b - 1.$$  \hspace{1cm} (A.10)

The inequality in relation (A.10) holds for $\chi, b \geq 0$. Equation (A.10) illustrates that for already moderately high $\chi$, potentially enhanced by a non-zero $b$, one would need rather high moments of the temperature to represent the effective excitation temperature correctly. The consideration does not allow to identify an optimal $n$ but nevertheless illustrates that $n = 4$ is not an unreasonable choice taking the excitation-like temperature as criterion.

Appendix B: Base RGB versus tip RGB abundance corrections

As pointed out above, the abundance corrections derived in the present work for the red giants located near the base of the RGB ($T_{\text{eff}} = 5000$ K, $\log g = 2.5$, $[\text{M/H}] = 0.0$, $-1.2$, $-3$) show little variation with wavelength, at least in the range 400–1600 nm. In contrast, Paper II demonstrated for a cooler red giant located near the tip of the RGB ($T_{\text{eff}} = 3600$ K, $\log g = 1.0$, $[\text{M/H}] = 0.0$) that the theoretical abundance corrections at 1600 nm were significantly smaller than those at optical wavelengths. In the following, we shall provide some basic explanation for the different wavelength dependence of the abundance corrections in these two type of giants.

As an example, we consider the formation of a weak fictitious high-excitation Fe II line with $\chi = 6$ eV, representative of the ionized atoms that show significant abundance corrections ([M/H] = 0.0). In the case of the tip RGB giant, the abundance correction derived for this line is $\Delta_{3D-3D} \approx -0.4$ dex at $\lambda = 850$ nm, and $\Delta_{3D-3D} \approx -0.02$ dex at $\lambda = 1600$ nm. In
The 3D abundance corrections are therefore expected to be po-
in the cool giant, and smallest in the warmer metal-poor giant. The metallicity dependence of the corrections is weak at $\lambda = 850$ nm and negligible at $\lambda = 1600$ nm (see Figs. 7 and 8). Since the corrections in the infrared are small for both type of gi-
ants, the remaining question is why the corrections in the red at $\lambda = 850$ nm are so much smaller in the “warm” giant than in the more evolved “cool” giant.

As in Paper II, the following considerations are restricted to vertical rays (disk-center intensity) in a single snapshot from the different 3D simulations. We focus here on the analysis of the “granulation correction”, $\Delta_{3D}(T,3D)$.

### B.1. Temperature fluctuations on iso-surfaces of optical depth

We define the mean height of line formation as the center of gravity of the equivalent width contribution function $B$ on the monochromatic continuum optical depth scale $\tau_c$:

$$\langle \log \tau_c \rangle = \frac{\int_{-\infty}^{\infty} \log \tau_c^* B(\tau_c^*) \text{d} \log \tau_c^*}{\int_{-\infty}^{\infty} B(\tau_c^*) \text{d} \log \tau_c^*}. \quad (B.1)$$

For simplicity, we use the contribution function of the (3D) atmosphere, corresponding to $B_{2,2,2}$ in Paper II (Eq. (B.10)). The results for the Fe\textsc{ii} line with $\chi = 6$ eV are given in Table B.1 for three different model atmospheres at $\lambda = 850$ and 1600 nm, respectively. The mean height of formation on the Rosseland optical depth scale, $\langle \log \tau_{Ross} \rangle$, is defined such that the average temperature on the iso-surface $\log \tau_{Ross} = \langle \log \tau_{Ross} \rangle$ is equal to the average temperature on the iso-surface $\log \tau_c = \langle \log \tau_c \rangle$, which we denote as $\langle T \rangle_{line}$. For all three models listed in Table B.1, the infrared line forms in significantly deeper atmospheric layers than the red line. This is the expected result since the continuum opacity is mainly due to $H^-$ and shows a local maximum near $\lambda$ 850 nm, and a minimum close to $\lambda$ 1600 nm. In the metal-poor giant, the line contribution function is more asymmetric and ex-
tends towards higher atmospheric layers than in the solar metal-
licity giant at the same $T_{eff}$ and log $g$. This is related to the fact that the continuum opacity decreases more steeply with height in the metal-poor atmosphere (due to the reduced electron pres-
sure). For given stellar parameters, the mean height of line for-
novation is therefore shifted towards higher atmospheric layers at low metallicity.

Table B.1 also gives the relative rms temperature fluctuation $\delta T_{rms}/\langle T \rangle$ versus the average temperature $\langle T \rangle$, evaluated on different iso-optical depth surfaces, $\tau_{Ross}(\lambda = 850)$ (solid), $\tau_{Ross}(\lambda = 1600)$ (dashed), $\tau_{Ross}$ (dotted). Filled circles indicate the values at the mean height of line formation at $\lambda 850$ nm and $\lambda 1600$ nm, respectively, as given in Table B.1.

### B.2. Opacity fluctuations and abundance corrections

We recall that, in the weak line limit, the abundance correction $\Delta_{3D}(T,3D)$ can be computed as

$$\Delta_{3D}(T,3D) = -\log \left( \frac{W_{3D}}{W_{3D}} \right), \quad (B.2)$$

where $W_{3D}$ and $W_{3D}$ denote the line equivalent width obtained from the 3D and the (3D) model, respectively, for the same atomic line parameters and elemental abundance (see Paper II, Appendix B). With the help of the equivalent width contribution function $B$, we can write the ratio of equivalent widths as

$$\frac{W_{3D}}{W_{3D}} = \frac{\int_{-\infty}^{\infty} B_{3,3,3}(\tau_c^*) \text{d} \log \tau_c^*}{\int_{-\infty}^{\infty} B_{2,2,2}(\tau_c^*) \text{d} \log \tau_c^*} \equiv \left( \begin{array}{c} W_{3,3,3} \\ W_{2,2,2} \end{array} \right) \equiv \mathcal{R}_{3,3,3}, \quad (B.3)$$

with the mixed contribution function

$$B_{i,j,k}(\tau_c) = \ln 10 \tau_c \exp[\tau_c] \left( \begin{array}{c} u_{c,j}(\tau_c) \\ \kappa_{c,j}(\tau_c) \end{array} \right) \left( \begin{array}{c} \kappa_{c,k}(\tau_c) \end{array} \right) \left( \begin{array}{c} \tau_c \end{array} \right) \quad (B.4)$$
Table B.1.  Mean height of formation, \((\log \tau_c)\), of a fictitious Fe \(\text{II}\) line with \(\chi = 6\text{ eV}\) at 850 and 1600 nm, respectively, for the red giant model discussed in Paper II, and of the CO\(^{18}\)BOLD red giant models used in this work.

| Model name           | \(T_{\text{eff}}\) [K] | \(\log g\) [cgs] | [M/H] | \(\langle \log \tau_c \rangle\) | \(\langle \log \tau_{\text{Ross}} \rangle\) | \(\langle (T)_{\text{line}} \rangle\) [K] | \(\delta T_{\text{rms}}\) [\%] | \(\tau_c\) |
|----------------------|-------------------------|------------------|-------|-----------------------------|-----------------------------------|-------------------------------------------|---------------------------------|------------|
|                      | \(\lambda\) 850 nm      |                  |       |                             |                                    |                                           |                                 |            |
| d3 t36 g10 mm00      | 3600                    | 1.0              | 0     | 0.53                        | 0.32                               | 4480                                      | 8.15                           | 6.33       |
| d3 t50 g25 mm00      | 5000                    | 2.5              | 0     | -0.01                       | -0.06                              | 5260                                      | 5.21                           | 4.68       |
| d3 t50 g25 mm20      | 5000                    | 2.5              | -2    | -0.36                       | -0.54                              | 4770                                      | 2.91                           | 2.87       |
|                      | \(\lambda\) 1600 nm     |                  |       |                             |                                    |                                           |                                 |            |
| d3 t36 g10 mm00      | 3600                    | 1.0              | 0     | -0.03                       | 0.68                               | 5120                                      | 6.48                           | 7.46       |
| d3 t50 g25 mm00      | 5000                    | 2.5              | 0     | -0.47                       | 0.15                               | 5600                                      | 4.31                           | 5.23       |
| d3 t50 g25 mm20      | 5000                    | 2.5              | -2    | -0.72                       | -0.16                              | 5160                                      | 4.00                           | 4.49       |

Notes. \((\log \tau_{\text{Ross}})\) is the mean height of formation on the Rosseland optical depth scale; \(\langle (T)_{\text{line}} \rangle\) is the average temperature at the mean height of formation. The last two columns give the relative rms temperature fluctuation on the iso-surface \(\log \tau_c = \langle \log \tau_c \rangle\) and \(\log \tau_{\text{Ross}} = \langle \log \tau_{\text{Ross}} \rangle\), respectively.

as defined in Paper II, Eq. (B.10). Subscripts 2 and 3 refer to the (3D) model and the 3D model, respectively.

To isolate the role of the line opacity fluctuations for the abundance corrections, we consider the ratio

\[
R_{2,3,2} \equiv \frac{W_{2,2,2}}{W_{2,2,2}} = \frac{\int_{-\infty}^{\infty} B_{2,2,2}(\tau_c') \, d \log \tau_c'}{\int_{-\infty}^{\infty} B_{2,2,2}(\tau_c') \, d \log \tau_c} = \frac{\int_{-\infty}^{\infty} B_{2,2,2}(\tau_c') \, A_{2,3,2}(\tau_c') \, d \log \tau_c'}{\int_{-\infty}^{\infty} B_{2,2,2}(\tau_c') \, d \log \tau_c'},
\]

where we have defined the local amplification factor \(A_{2,3,2}\) as

\[
A_{2,3,2}(\tau_c) = \frac{B_{2,3,2}(\tau_c)}{B_{2,2,2}(\tau_c)} = \frac{\eta_{c,3}(\tau_c)}{\eta_{c,2}(\tau_c)} = \frac{(\eta(T))_{\text{g}y}}{\eta(T)_{\text{g}y}},
\]

According to Eq. (B.5), the equivalent width ratio \(R_{2,3,2}\) can be expressed as an average of the amplification factor \(A_{2,3,2}\) over optical depth with weighting function \(B_{2,2,2}\). Figure B.2 (left column) shows the depth dependence of \(A_{2,3,2}\) at both wavelengths for the three red giant models under consideration, together with the weighting function \(B_{2,2,2}\) and the resulting values of \(R_{2,3,2}\). The largest 3D effects are found in the cool giant. The reason is twofold: (i) the amplitude of the temperature fluctuations in the line forming layers of the cool giant is significantly higher than in the warmer giants (see above); (ii) the Fe \(\text{II}\) ionization fraction is strongly variable in the atmosphere of the cool giant, especially in the layers where \(\lambda 850\) nm line forms, while Fe \(\text{II}\) is always the dominating ionization stage in the warmer giants. Both factors enhance the nonlinearity of the line opacity fluctuations in the cool giant, and lead to the pronounced 3D abundance corrections.

Next we consider the combined effect of line and continuum opacity fluctuations. For this purpose, we have to evaluate the amplification factor \(A_{2,3,3}\):

\[
A_{2,3,3}(\tau_c) = \frac{B_{2,3,3}(\tau_c)}{B_{2,2,2}(\tau_c)} = \frac{(\eta_{c,3}(\tau_c))_{\text{g}y}}{\eta_{c,2}(\tau_c)} = \frac{(\eta(T))_{\text{g}y}}{\eta(T)_{\text{g}y}},
\]

where \(\eta_{c} = \kappa_{c}/\kappa_{f}\) is the ratio of line to continuum opacity. Again, the effective amplification factor \(R_{2,3,3}\) is obtained as the average of the local amplification factor \(A_{2,3,3}\) over optical depth with weighting function \(B_{2,2,2}\). The result is shown in Fig. B.2 (right column). Obviously, the line and the continuum opacity are correlated in the sense that both increase with temperature, such that the fluctuation amplitude of their ratio, \(\eta_{c}\), is always lower than the fluctuation amplitude of the line opacity, \(\kappa_{c}\). Hence, \(R_{2,3,3}\) is smaller than \(R_{2,3,2}\). For the warmer giants, the reduction of \(R\) is of similar magnitude for both wavelengths, \(\log(R_{2,3,3}/R_{2,3,2}) = -0.05\) dex. For the cool giant, however, the reduction factor is much smaller at \(\lambda 850\) nm (\(=0.01\) dex) than at \(\lambda 1600\) nm (\(=0.11\) dex), further increasing the difference between the 3D corrections derived for the two wavelengths.

The similarity of \(R_{2,3,2}\) and \(R_{2,3,3}\) for the \(\lambda 850\) nm line in the cool giant is implied by the similarity of the local amplification factors \(A_{2,3,2}\) and \(A_{2,3,3}\) in the line forming region. The reason for this behavior can be traced back to the low temperature sensitivity of the continuum opacity in the temperature range 4000 K \(< T < 5000\) K (cf. Fig. B.2, in Paper II), which coincides with the formation region of this line (see blue weighting function in the upper panels of Fig. B.2). For the warmer giant with solar metallicity, a similar “plateau” in the \(\kappa_{c}\) versus \(T\) relation is found in the same temperature range. However, in this giant the \(\lambda 850\) nm line forms at higher temperatures (see blue weighting function in the middle panels of Fig. B.2), and the temperature sensitivity of \(\kappa_{c}\) is thus comparable at both wavelengths. Remarkably, the “plateau” in the \(\kappa_{c}\) versus \(T\) relation is not seen in the metal-poor giants. Presumably, the “plateau” is related to the transition from a hydrogen to a metal dominated electron source, the latter being negligible in a metal-poor atmosphere.

Finally, we have also evaluated the full 3D corrections \(R_{3,3,3}\), marked by star symbols in Fig. B.2 (right panels). The additional effect of the fluctuations of the source function gradient \(u_{c}\) is only significant in the cool giant, and increases the dichotomy between the corrections at \(\lambda 850\) and 1600 nm even further. A similar behavior was found for the Fe \(\text{II}\) line with \(\chi = 10\text{ eV}\) investigated in Paper II.

In summary, the 3D corrections for high-excitation lines of ions at wavelengths near \(\lambda 850\) nm are significantly larger in the cool tip RGB giant discussed in Paper II than in the warmer giants near the base of the RGB studied in the present work because, in the cool giant, the lines form in a region where (i) the temperature fluctuations are larger, (ii) the temperature sensitivity of the continuum opacity is particularly low, and (iii) the ionization fraction of typical ions like Fe \(\text{II}\) varies considerably, thus enhancing the temperature sensitivity of the line opacity.
Appendix C: Formation of molecular lines

Horizontal temperature fluctuations play an essential role in defining abundance corrections for the molecules. Firstly, they are larger in the 3D hydrodynamical models at lower metallicity (cf. Figs. 1, 2), which leads to the highest molecular number density and line opacity fluctuations in the lowest metallicity models. Secondly, in a model of a given metallicity different molecules form at different optical depths which are characterized by different amplitude of the horizontal temperature fluctuations. For example, CO formation occupies the outermost range in terms of Rosseland optical depth of all molecules studied here (Fig. C.2). Consequently, its lines form in the atmospheric layers characterized by the largest horizontal temperature fluctuations (cf. Fig. 2) in comparison to those in the layers where other molecules form. The fluctuations of the line opacity is therefore largest in the case of CO in the 3D model at [M/H] = −3.0, which is part of the explanation of the large abundance corrections for CO. However, further factors are important.

According to Paper II, the number density of molecules per unit mass (calculated as $X_i = N_i/\rho$, here $\rho$ is the mass density) can be computed using the following Saha-like equation

$$X_{AB} = \frac{\rho}{\rho_0} \frac{Q_{AB}(T)}{U_A(T) U_B(T)} \frac{X_B}{X_A} \left( \frac{h^2}{2\pi m_{AB} k T} \right)^{3/2} e^{-D_{0}/kT}. \ (C.1)$$

where $Q_{AB}$ is the partition function of the molecule AB, $X_A$ and $X_B$ are number densities per unit mass of the constituent species A and B, with their partition functions $U_A(T)$ and $U_B(T)$, respectively, mass of the molecule $m_{AB}$, its dissociation energy $D_0$, and temperature, $T$. The number densities of the molecules studied here are shown in Fig. C.1 plotted versus Rosseland optical depth in the 1D LHD model at [M/H] = −3.0 (number densities are normalized to the total number density of carbon nuclei, $X_{AB}/\sum X_C$). Qualitatively, the situation is very similar to that in
Fig. C.1. Number density of different molecules, normalized to the total number density of carbon, plotted as a function of the Rosseland optical depth in the 1D LHD model at $[M/H] = -3.0$.

the cool red giant studied by Paper II: because the abundance of oxygen is higher than that of carbon, most of the available C is locked into CO beyond $\log \tau_{Ross} \sim -0.5$. In the deeper atmosphere, the molecules are quickly destroyed due to dissociation.

From Eq. (C.1) one can expect rather different temperature sensitivity of the number densities of different molecules. Indeed, this is clearly seen in Fig. C.3 where we show number (normalized) densities of CO, CN, CH, C$_2$, NH and OH in the 3D model ([M/H] = -3.0) at different levels of constant Rosseland optical depths, plotted versus inverse temperature $\theta = 5040/T$. It is obvious that CO is significantly more sensitive to changes in $T$ than the other molecules, essentially due to the high dissociation energy of CO, $D_0 = 11.1$ eV. In fact, the temperature sensitivity of the molecular number densities scales with $D_0$ in the hotter parts of the atmosphere where $T \sim 4000$ K. In the cooler regions, however, where almost all carbon is locked up in CO and atomic carbon is a minority species, the temperature dependence of the number density of the other carbon-bearing molecules is reversed: here a temperature increase leads to the destruction of CO, and in turn, to a larger concentration of atomic carbon that enables an enhanced formation of carbon-bearing molecules. Carbon-free molecules do not show this anomalous behavior at low temperature (see lower panels of Fig. C.3).

Since the line opacity is directly proportional to $X_{AB}$, larger fluctuations of the number densities should lead to larger fluctuations and larger nonlinearities in the line opacities, and thus to larger abundance corrections. This is basically what one sees in case of CO: the broadest range of fluctuations in the case of CO leads to the largest abundance corrections amongst all molecules investigated here.

Interestingly, in the case of cool red giant studied in Paper II, the abundance corrections were largest for C$_2$. In contrast, the abundance corrections for C$_2$ obtained in this work are in fact smallest amongst all molecules. A closer look at the dependence of C$_2$ number density on temperature reveals that $X_{C2}$ increases steeply up to $T \sim 4000$ K and then decreases towards the higher temperatures (Fig. C.3, middle right). This is because the formation of C$_2$ is tightly coupled to that of CO, as explained above, and makes the number density of C$_2$ extremely sensitive to changes in temperature at low $T$. In the cool red giant analyzed by Paper II the atmosphere is significantly cooler and the C$_2$ line forms entirely the lower temperature regime where $X(C_2)$ is highly sensitive to the temperature fluctuations, leading to the largest abundance corrections for C$_2$. In the warmer red giant studied here, however, the C$_2$ line forms in the transition region between both temperature regimes, such that the amplitude of the fluctuations is limited and their net effect on the abundance corrections is much smaller.

NH ($D_0 = 3.5$ eV) and OH ($D_0 = 4.4$ eV) have similar molecular properties and line formation regions. The 3D abundance correction is slightly more negative for OH because this molecule has a somewhat higher dissociation energy. The correction for CN ($D_0 = 7.8$ eV) falls in the same range, although this molecule has a much higher dissociation energy. Again, the explanation for this unexpected behavior is related to the change of sign in the temperature sensitivity of the number density of CN in the line forming region ($\log \tau_{Ross} = -2$), due to the coupling with CO at low temperatures. The coupling with CO is missing in the case of carbon-free molecules NH and OH, where the slope $\partial \log N/\partial \theta$ is therefore essentially constant over the whole temperature range (see Fig. C.3, lower panels).
Fig. C.2. Equivalent width contribution function (flux) of weak ($W \leq 0.5$ pm) artificial molecular lines of CO, CN, CH, C$_2$, NH, OH (top left to bottom right) with lower level excitation potential $\chi = 0$ eV in the full 3D, average (3D), and 1D model atmospheres with metallicity [M/H] = −3.0. Contribution functions were normalized to the equivalent width of the line computed using 3D models.
Fig. C.3. Normalized number densities of molecules CO, CN, CH, C$_2$, NH, OH (top left to bottom right) in the 3D model atmosphere ([M/H] = −3.0) plotted at different levels of constant Rosseland optical depth (identified by color) as a function of inverse temperature $\theta = 5040/T$. 