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ABSTRACT

The calculation of the thermal stratification in the superadiabatic layers of stellar models with convective envelopes is a long standing problem of stellar astrophysics, and has a major impact on predicted observational properties like radius and effective temperature. The Mixing Length Theory, almost universally used to model the superadiabatic convective layers, contains effectively one free parameter to be calibrated $-\alpha_{ml}$ whose value controls the resulting effective temperature. Here we present the first self-consistent stellar evolution models calculated by employing the atmospheric temperature stratification, Rosseland opacities, and calibrated variable α_{ml} (dependent on effective temperature and surface gravity) from a large suite of three-dimensional radiation hydrodynamics simulations of stellar convective envelopes and atmospheres for solar stellar composition (Trampedach et al. 2013). From our calculations (with the same composition of the radiation hydrodynamics simulations), we find that the effective temperatures of models with the hydrocalibrated variable α_{ml} (that ranges between ~1.6 and ~2.0 in the parameter space covered by the simulations) display only minor differences, by at most ~30-50 K, compared to models calculated at constant solar α_{ml} (equal to 1.76, as obtained from the same simulations). The depth of the convective regions is essentially the same in both cases. We have also analyzed the role played by the hydro-calibrated $T(\tau)$ relationships in determining the evolution of the model effective temperatures, when compared to alternative $T(\tau)$ relationships often used in stellar model computations. The choice of the $T(\tau)$ can have a larger impact than the use of a variable α_{ml} compared to a constant solar value. We found that the solar semi-empirical $T(\tau)$ by Vernazza et al. (1981) provides stellar model effective temperatures that agree quite well with the results with the hydro-calibrated relationships.

Key words. convection - stars: atmospheres - stars: evolution - stars: Hertzsprung-Russell and C-M diagrams

1. Introduction

The almost universally adopted method for calculating superadiabatic convective temperature gradients in stellar evolution models is based on the formalism provided by the so-called Mixing Length Theory (MLT - Böhm-Vitense 1958). This formalism is extremely simple; the gas flow is made of columns of upward and downward moving convective elements with a characteristic size, the same in all dimensions, that cover a fixed mean free path before dissolving. All convective elements have the same physical properties at a given distance from the star centre; upward moving elements release their excess heat into the surrounding gas, and are replaced at their starting point by the downward moving elements, that thermalize with the surrounding matter, thus perpetuating the cycle. The MLT is a 'local' theory, and the evaluation of all relevant physical and chemical quantities are based on the local properties of each specific stellar layer, regardless of the extension of the whole convective region.

Both the mean free path and the characteristic size of the convective elements are assumed to be same for all convective bubbles, and are assigned the same value $\Lambda = \alpha_{ml}H_p$, the so-called 'mixing length'). Here α_{ml} is a free parameter (assumed to be a constant value within the convective regions and along all evolutionary phases), and H_p is the local pressure scale height. There are additional free parameters in the MLT, that are generally fixed a priori (versions of the MLT with different choices of these additional parameters will be denoted here as different statement.

ent MLT 'flavours'), so that practically the only free parameter to be calibrated is α_{ml} . Stellar evolution calculations for a fixed mass and initial chemical composition but with varying α_{ml} , produce evolutionary tracks with different T_{eff} (and radius) evolution, whereas evolutionary timescales and luminosities are typically unchanged. On the other hand, the prediction of accurate values of T_{eff} (and radii) by evolutionary models and stellar isochrones is paramount, among others, to study colourmagnitude diagrams of resolved stellar populations, empirical mass-radius relations of eclipsing binary systems, and predict reliable integrated spectra (and colours) of unresolved stellar systems (extragalactic clusters and galaxies).

In stellar evolution calculations the value of α_{ml} is usually calibrated by reproducing the radius of the Sun at the solar age with an evolutionary solar model. This solar calibrated α_{ml} is then kept fixed in all evolutionary calculations of stars of different masses and chemical compositions. The exact numerical value of α_{ml} varies amongst calculations by different authors because variations of input physics and choices of the outer boundary conditions affect the predicted model radii and T_{eff} values, hence require different α_{ml} values to match the Sun. Regarding the various MLT flavours, Pedersen et al. (1990) and Salaris & Cassisi (2008) have shown how they provide the same T_{eff} evolution, once α_{ml} is appropriately recalibrated on the Sun.

As discussed by, e.g., Vandenberg et al. (1996) and Salaris et al. (2002) metal poor red giant branch (RGB) models calculated with the solar calibrated value of α_{ml} (hereafter

 $\alpha_{ml,\odot}$) are able to reproduce the $T_{\rm eff}$ of samples of RGB stars in Galactic globular clusters, within the error bars on the estimated $T_{\rm eff}$. On the other hand, several authors find that variations of $\alpha_{\rm ml}$ with respect to $\alpha_{\rm ml,\odot}$ are necessary to reproduce –just to give some examples– the red edge of the RGB in a sample of 38 nearby Galactic disc stars with radii determined from interferometry (Piau et al. 2011), the asteroseismically constrained radii of a sample of main sequence (MS) *Kepler* targets (Mathur et al. 2012), observations of binary MS stars in the Hyades (Yıldız et al. 2006) and the α Cen system (Yıldız 2007), a low-mass pre-MS eclipsing binary in Orion (Stassun et al. 2004).

The MLT formalism provides only a very simplified description of convection, and there have been several attempts to introduce non-locality in the MLT (see, e.g., Grossman et al. 1993; Deng et al. 2006, and references therein). These 'refinements' of the MLT are often complex and introduce additional free parameters to be calibrated. The alternative model by Canuto & Mazzitelli (1991) and Canuto & Mazzitelli (1992) includes a spectrum of eddy sizes (rather than the one-sized convective cells of the MLT) and fixes the scale length of the convective motions to the distance to the closest convective boundary. Recently Pasetto et al. (2014) have presented a new non-local and time-dependent model based on the solution of the Navier-Stokes equations for an incompressible perfect fluid, that does not contain any free parameter¹.

An alternative approach to model the superadiabatic layers of convective envelopes is based on the computation of realistic multidimensional radiation hydrodynamics (RHD) simulations of atmospheres and convective envelopes -where convection emerges from first principles- that cover the range of effective temperatures (T_{eff}) , surface gravities (g), and compositions typical of stars with surface convection. These simulations have reached nowadays a high level of sophistication (see, e.g., Nordlund et al. 2009) and for ease of implementation in stellar evolution codes, their results can be used to provide an 'effective' hydro-calibration of α_{ml} , even though RHD simulations do not confirm the basic MLT picture of columns of convective cells. After early attempts from rather crude twodimensional (2D) and three-dimensional (3D) simulations (see, e.g., Deupree & Varner 1980; Lydon et al. 1992), a first comprehensive RHD calibration of α_{ml} was presented by Ludwig et al. (1999) and Freytag et al. (1999). These authors found from their simulations that the calibrated α_{ml} varies as a function of metallicity, g and $T_{\rm eff}$. Freytag & Salaris (1999) applied this RHD calibration of $\alpha_{\rm ml}$ (plus $T(\tau)$ relations computed from the same RHD models, and Rosseland opacities consistent with the RHD calculations) to metal poor stellar evolution models for Galactic globular cluster stars, and found that the resulting isochrones for the relevant age range have only small $T_{\rm eff}$ differences along the RGB (of the order of ~50 K) with respect to isochrones computed with a solar calibrated value of α_{ml} .

In the last years a number of grids of 3D hydrodynamics simulations of surface convection have been published, and from the point of view of stellar model calculations it is very important to study whether the results by Freytag & Salaris (1999) are confirmed or drastically changed.

Tanner et al. (2013b) and Tanner et al. (2013a) have presented a grid of simulations employing in the optically thin layers a 3D Eddington solver (Tanner et al. 2012). Their calculations cover four metallicities (from Z=0.001 to Z=0.04), but just a few $T_{\rm eff}$ values at constant $\log(g)$ =4.30 for each Z, plus a subset of models at varying He mass fraction for Z=0.001 and Z=0.02. These authors studied the properties of convection with varying $T_{\rm eff}$ and chemical composition in these solar-like envelopes. Tanner et al. (2014) extracted metallicity-dependent $T(\tau)$ relations from these same simulations, and employed them to highlight the critical role these relations play when calibrating $\alpha_{\rm ml}$ with stellar evolution models.

Magic et al. (2013) have published a very large grid of 3D RHD simulations for a range of chemical compositions. Their grid covers a range of $T_{\rm eff}$ from 4000 to 7000 K in steps of 500 K, a range of log(g) from 1.5 to 5.0 in steps of 0.5 dex, and metallicity, [Fe/H], from -4.0 to +0.5 in steps of 0.5 and 1.0 dex. These models have been employed by Magic et al. (2015) to calibrate $\alpha_{\rm ml}$ as function of g, $T_{\rm eff}$ and [Fe/H]. They found that $\alpha_{\rm ml}$ depends in a complex way on these three parameters, but in general $\alpha_{\rm ml}$ decreases towards higher effective temperature, lower surface gravity and higher metallicity. So far Magic et al. (2015) have provided only fitting formulae for $\alpha_{\rm ml}$ but not publicly available prescriptions for the boundary conditions and input physics.

Very recently Trampedach et al. (2013) produced a nonsquare grid of convective atmosphere/envelope 3D RHD simulations for the solar chemical composition. The grid spans a $T_{\rm eff}$ range from 4200 to 6900 K for MS stars around $\log(g)=4.5$, and from 4300 to 5000 K for red giants with log(g)=2.2, the lowest surface gravity available. In Trampedach et al. (2014b) the horizontal and temporal averages of the 3D simulations were then matched to 1D hydrostatic equilibrium, spherically symmetric envelope models to calibrate α_{ml} as function of g and $T_{\rm eff}$ (see Trampedach et al. 2014b, for details about the calibration procedure). Moreover, the same RHD simulations have been employed by Trampedach et al. (2014a) to calculate g- and $T_{\rm eff}$ -dependent $T(\tau)$ relations from temporal and τ (Rosseland optical depth) averaged temperatures of the atmospheric layers. Trampedach et al. (2014b) also provide routines to calculate their g- and $T_{\rm eff}$ -dependent RHD-calibrated $\alpha_{\rm ml}$ together with their computed $T(\tau)$ relations, and Rosseland opacities consistent with the opacities used in the RHD simulations. This enables stellar evolution calculations where boundary conditions, superadiabatic temperature gradient and opacities of the convective envelope are consistent with the RHD simulations. It is particularly important to use both the RHD-calibrated α_{ml} and $T(\tau)$ relations, because the $T_{\rm eff}$ of the stellar evolution calculations depends on both these inputs (see, i.e., Salaris et al. 2002; Tanner et al. 2014, and references therein).

Thanks to this consistency between RHD simulations and publicly available stellar model inputs, we present and discuss in this paper the first stellar evolution calculations where Trampedach et al. (2014b) 3D RHD-calibration of α_{ml} is selfconsistently included in the evolutionary code. In the same vein as Freytag & Salaris (1999), we focus on the effect of the calibrated variable α_{ml} on the model T_{eff} , compared to the case of calculations with fixed $\alpha_{ml,\odot}$ (as determined from the same RHD simulations). Self consistency of opacity and boundary conditions is paramount to assess correctly differential effects, given that the response of models to variations of α_{ml} depends on their T_{eff} , that in turn depends on the absolute value of α_{ml} , opacities and boundary conditions. We also address the role played by the RHD calibrated $T(\tau)$ relations in the determination of the model T_{eff} when compared to other widely used relations.

Section 2 describes briefly the relevant input physics of the models, while Sect. 3 presents and compares the resulting evolutionary tracks. A summary and discussion close the paper.

¹ This model has not been implemented yet in any stellar evolution computations.

2. Input physics

All stellar evolution calculations presented here have been performed with the BaSTI (a BAg of Stellar Tracks and Isochrones) code (Pietrinferni et al. 2004), for a chemical composition with Y=0.245, Z=0.018 and the Grevesse & Noels (1993) metal mixture, consistent with the chemical composition of the RHD simulations. Atomic diffusion was switched off in these calculations, and convective core overshooting was included when appropriate (see Pietrinferni et al. 2004, for details).

We employed the Rosseland opacities provided by Trampedach et al. (2014b). The low temperature opacities (for $\log(T)<4.5$) are very close to Ferguson et al. (2005) with the exception of the region with $\log(T)<3.5$, where Ferguson et al. (2005) calculations are higher because of the inclusion of the effect of water molecules (see Trampedach et al. 2014a). Opacity Project (Badnell et al. 2005) calculations are used for $\log(T)\geq4.5$. As for the surface boundary conditions, we employed the $T(\tau)$ relations computed by Trampedach et al. (2014a), who provided a routine that calculates, for a given T_{eff} and surface gravity, the appropriate generalized Hopf functions $q(\tau)$, related to the $T(\tau)$ relation by

$$q(\tau) = \frac{4}{3} \left(\frac{T(\tau)}{T_{\text{eff}}} \right)^4 - \tau \tag{1}$$

In our calculations we fixed to $\tau_{tr}=2/3$ the transition from the $T(\tau)$ integration of the atmospheric layers (with τ as independent variable) to the integration of the full system of stellar structure equations. As mentioned by Trampedach et al. (2014a) and Trampedach et al. (2014b), these RHD-based $T(\tau)$ relations can be employed to model also the convective layers in the optically thick part of the envelope, together with modified expressions for the temperature gradients in the superadiabatic regions, and an appropriate rescaling of τ (see Trampedach et al. 2014a, for details). In our calculations we have compared the radiative (∇_{rad}) and superadiabatic (∇ –obtained with the RHD calibrated values of α_{ml}) temperature gradients determined along the upper part of the convective envelopes from the standard stellar structure equations and MLT (down to τ =100, the upper limit for the routine calculating the Hopf functions), with ∇_{rad} and ∇ calculated according to Eqs.35 and 36 of Trampedach et al. (2014a), respectively. We have found that in all our models the differences between these two sets of gradients are much less than 1% between $\tau_{\rm tr}$ and $\tau = 100$.

We have calculated also test models by changing $\tau_{\rm tr}$ between 2/3 and 5 (with the appropriate rescaling of τ if convection appears in the atmosphere integration, see Trampedach et al. 2014a), and obtained identical evolutionary tracks in each case.

Regarding the value of α_{ml} , the same routine for the Hopf functions provides also the RHD calibrated value of α_{ml} (for the Böhm-Vitense 1958, flavour of the MLT) for a given T_{eff} and surface gravity, that we employed in our calculations. Uncertainties in the calibrated α_{ml} values are of the order of $\pm 0.02 - 0.03$ (see Table 1 of Trampedach et al. 2014b)².

The only difference in terms of input physics between the atmosphere/envelope RHD calculations and our models is the equation of state (EOS). The RHD simulations employed the MHD (Daeppen et al. 1988) EOS, that is not the same EOS used in the BaSTI calculations (see Pietrinferni et al. 2004). To check whether this can cause major differences in the models, we have calculated envelope models for the same g- $T_{\rm eff}$ pairs of the RHD simulations, including the RHD-calibrated $\alpha_{\rm ml}$, $T(\tau)$ relations, and RHD opacities. We have then compared the resulting depths of the convection zones ($d_{\rm CZ}$, in units of stellar radius) with what obtained by Trampedach et al. (2014b) from their RHD-calibrated 1D envelope models, that used the same input physics (including EOS) of the RHD calculations (see Table 1 of Trampedach et al. 2014b). We found random (non systematic) differences of $d_{\rm CZ}$ by at most just 2-3% compared to Trampedach et al. (2014b) results.

3. Model comparisons

As mentioned in the introduction, the RHD simulations cover a non-square region in the $g-T_{eff}$ diagram, as displayed in Fig. 1 (the region enclosed by thick solid lines), ranging from 4200 to 6900 K on the MS, and from 4300 to 5000 K for RGB stars with $\log(g)=2.2$. The MLT calibration results in an α_{ml} varying from 1.6 for the warmest dwarfs, with a thin convective envelope, up to 2.05 for the coolest dwarfs in the grid. In between there is a triangular plateau of $\alpha_{ml} \sim 1.76$, where the Sun is located. The RHD simulation for the Sun provides $\alpha_{ml} \sim 1.76\pm0.03$. The top panel of Fig. 1 displays the results of our evolutionary model calculations in the $g-T_{eff}$ diagram (from the pre-MS to the lower RGB), for masses M=0.75, 1.0, 1.4, 2.0 and $3.0M_{\odot}$ respectively, that cover the full domain of the RHD simulations.

The thick solid lines denote the reference set of models, calculated with the varying α_{ml} calibration, and the run of α_{ml} along each individual track is displayed in the lower panel. The dotted part of each sequence in this panel denotes the α_{ml} values extrapolated by the calibration routine, when the models are outside the region covered by the simulations but still retain a convective region. This happens along the pre-MS evolution of the M=0.85M_o track, and for the 2.0 and 3.0M_o calculations along the subgiant phase. Apart from the pre-MS stages of the two lowest mass models, the evolution of α_{ml} spans a narrow range of values, between ~1.6 and ~1.8.

To study the significance of the variation of α_{ml} for the model T_{eff} , we have calculated evolutionary models for the same masses, this time keeping $\alpha_{ml} = \alpha_{ml,\odot} = 1.76$ along the whole evolution. The results are also displayed in Fig. 1.

A comparison of the two sets of tracks clearly shows that the effect of a varying α_{ml} is almost negligible. The largest differences are of only ~30 K along the RGB phase of the 3.0M_o track (solar α tracks being hotter because of a higher α_{ml} value compared to the calibration) and ~50 K at the bottom of the Hayashi track of the 1.0M_o track (solar α_{ml} tracks being cooler, because of a lower α_{ml}). In all other cases differences are smaller, and often equal to almost zero.

For all stellar masses we found that the mass fraction of He dredged to the surface by the first dredge up –that depends on the maximum depth of the convective envelope at the beginning of the RGB phase– is the same within 0.001, between constant $\alpha_{ml,\odot}$ and variable α_{ml} models. We have then compared the luminosity of the RGB bump –that also depends on the maximum depth of the convective envelope at the first dredge up (see, e.g., Cassisi & Salaris 1997, 2013, and references therein)– for the 0.75 and 1.0M_{\odot} models. We found that the luminosity is unchanged between models with constant and variable α_{ml} . This reflect the fact that the depth of the convective envelope is the same between the two sets of models throughout the MS phase to the RGB, until the end of the calculations. Small differences appear

² We did not include any turbulent pressure in the convective envelope, as the α_{ml} RHD calibration was performed in a way that works for standard stellar evolution models without this extra contribution to the pressure (R. Trampedach private communication, see also Sect.4 from Trampedach et al. 2014b)



Fig. 1. Stellar evolution tracks in the g- T_{eff} diagram for the labelled masses. The region enclosed by the thick black boundary is the g- T_{eff} range covered by the RHD simulations. Thick solid lines denote fully consistent calculations with the RHD calibrated variable α_{ml} and $T(\tau)$ relationships. The lower panel displays the evolution of α_{ml} along each track. The dotted portion of each sequence denotes the region where the α_{ml} values are extrapolated. Dashed lines in the upper panel display tracks calculated with a constant $\alpha_{\text{ml}} = \alpha_{ml,\odot}$ and the calibrated $T(\tau)$ relationships.

only during the pre-MS. To this purpose, for the $1.0M_{\odot}$ models we have additionally checked the surface Li abundance that survives the pre-MS depletion. This provides information about the evolution of the lower boundary of the convective envelope during this phase, where according to Fig. 1, α_{ml} shows the largest difference from $\alpha_{ml,\odot}$. We found that models with constant $\alpha_{ml,\odot}$ display after the pre-MS a Li abundance just 9% higher than the reference results.

We have then analyzed the role played by the $T(\tau)$ relationships computed from the RHD calculations in determining the model $T_{\rm eff}$ (the role played by the boundary conditions in determining the $T_{\rm eff}$ of stellar models is especially crucial for verylow-mass stars, see i.e. Allard et al. 1997; Brocato et al. 1998). Figure 2 displays the ratio T/T_{eff} as a function of τ as predicted by the calibrated relationships for atmospheres/envelopes with T_{eff} =4500 K and 6000 K (log(g)=3.5). The RHD calibrated $T(\tau)$ relationships contain values for the Hopf function that vary with τ and $T_{\rm eff}$ (also with g, to a lesser degree). The variation with $T_{\rm eff}$ is obvious from the figure. For these two temperatures the largest differences appear at the layers with τ between ~0.1 and \sim -1, where stellar model calculations usually fix the transition from the atmosphere to the interior. The same Figure 2 also displays the results for the traditional Eddington approximation to the grey atmosphere, and the solar semi-empirical $T(\tau)$ relationships by Krishna-Swamy (KS - Krishna Swamy 1966) and Vernazza et al. (1981) -their Model C for the quiet sun, hereinafter VAL_cT(τ). In these latter cases the ratio T/T_{eff} does not depend on $T_{\rm eff}$. It is easy to notice that around the photospheric



Fig. 2. Comparison of the ratio T/T_{eff} as a function of the optical depth τ as predicted by different $T(\tau)$ relationships. The thick solid and dotted lines display the RHD-calibrated relationships for T_{eff} =4500 K and 6000 K (log(g)=3.5 in both cases), respectively. The thin solid, dash-dotted and dashed lines show the ratios obtained from the Eddington, KS and VAL_c $T(\tau)$ relationships, respectively.

layers the RHD relationships are in between the Eddington grey and VAL_cT(τ). The most discrepant relationship is the KS one.

Figure 3 displays the results of evolutionary calculations with this set of $T(\tau)$ relationships. We compare here the reference set of models for 1.0 and $1.4 M_{\odot}$ and varying $\alpha_{\rm ml}$, with constant $\alpha_{ml,\odot}$ models calculated using the Eddington, KS and VAL_c $T(\tau)$ relationships (we set $\tau_{tr}=2/3$ also for these calculations).

Differences between these new sets of models at constant $\alpha_{ml,\odot}$ and the reference calculations are larger than the case of Fig. 1 because of the differences with the RHD $T(\tau)$ relationships. On the whole, the VAL_c $T(\tau)$ (coupled to the RHD calibrated $\alpha_{ml,\odot}$) gives the closest match to the self-consistent reference calculations. The largest differences appear for the $1.0M_{\odot}$ along the pre-MS; models calculated with the VAL_c relation are ~50 K cooler, approximately the same as the case of $\alpha_{ml,\odot}$ and the RHD $T(\tau)$. For the same $1.0M_{\odot}$ track the $T_{\rm eff}$ differences along the RGB are at most equal to 10 K, and at most ~40 K along the MS. Differences for the $1.4M_{\odot}$ track are smaller.

As for the Eddington T(τ), the resulting tracks are generally hotter than the self-consistent RHD-based calculations. The largest differences amount to ~40-50 K along MS and RGB of the two tracks. Comparisons with Fig. 1 show that, from the point of view of the resulting model T_{eff} , the Eddington $T(\tau)$ differs more –albeit by not much– than the VAL_c one from the RHD-calibrated relations.

The worse agreement is found with the KS T(τ). For both masses the RGB is systematically cooler by ~70 K, and the pre-MS by ~80-100 K, whilst the MS of the 1.0M_{\odot} calculations is cooler by ~120 K, a difference reduced to ~50 K along the MS of the 1.4M_{\odot} track.



Fig. 3. As the upper panel of Fig. 1 but for evolutionary tracks with masses equal to 1.0 and $1.4M_{\odot}$ respectively. Thick solid lines denote fully consistent calculations with the RHD calibrated variable α_{ml} and $T(\tau)$ relationships. Dotted, dash-dotted and dashed lines display tracks calculated with constant $\alpha_{ml,\odot}$ and the Eddington, KS and VAL_c T(τ), respectively.

In case of the $1.0M_{\odot}$ models we checked again the luminosity level of the RGB bump, the surface He mass fraction after the first dredge up and the amount of surface Li after the pre-MS depletion. For all these three $T(\tau)$ relations the evolutionary models display a RGB bump luminosity within $\Delta(L/L_{\odot}) < 0.01$, and a post-dredge up He abundance within 0.001 of the fully consistent result. In all these calculations the convective envelope has the same depth throughout the MS phase and along the RGB. Differences along the pre-MS are highlighted by the amount of Li depletion during this phase. We found that models calculated with the Eddington $T(\tau)$ have 9% less Li after the pre-MS, compared to the reference calculations. Comparing this number with just the effect of a constant $\alpha_{ml,\odot}$ discussed above, we derive that the use of this $T(\tau)$ decreases the surface Li abundance by ~20% compared to the use of the Hopf functions determined from the RHD simulations. In case of the VAL_c $T(\tau)$, the net effect is to increase the surface Li after the pre-MS by ~10% compared to the RHD T(τ), and also the KS relation causes a similar increase by ~11%.

3.1. The standard solar model

We close our analysis by discussing the implications of the RHD results and the choice of the $T(\tau)$ relation, on the calibration of the standard solar model. As well known, in stellar evolution it is customary to fix the value of $\alpha_{ml,\odot}$ (and the initial solar He and metal mass fractions) by calculating a $1M_{\odot}$ stellar model that matches the solar bolometric luminosity and radius at the age of the Sun, with the additional constraint of reproducing the present metal to hydrogen mass fraction Z/X ratio (see, e.g., Pietrinferni et al. 2004, for details). The accuracy of the derived solar model can then be tested against helioseismic estimates of

the depth of the convective envelope and the surface He mass fraction. It is also well established that solar models without microscopic diffusion cannot properly account for some helioseismic constraints, hence solar models are routinely calculated by including microscopic diffusion of He and metals (see, e.g., Pietrinferni et al. 2004, for a discussion and references).

We have first calculated a standard solar model (with the same input physics and solar metal distribution of the calculations discussed above) employing both the variable α_{ml} and the $T(\tau)$ relations from the RHD results. Given that microscopic diffusion decreases with time the surface chemical abundances of the model, the initial solar *Z* (and *Y*) need to be higher than the present one. This means that we had to employ the RHD results also for chemical compositions not exactly the same as the composition of the RHD simulations.

We found that it is necessary to rescale the RHD α_{ml} calibration by a factor of just 1.034 to reproduce the solar radius. This implies $\alpha_{ml,\odot}=1.82$, extremely close, within the error, to the RHD value $\alpha_{ml,\odot}=1.76\pm0.01$ (range) ±0.03 (calibration uncertainty) obtained by (Trampedach et al. 2014b). Given our previous results, it is also obvious that a solar model calibration with fixed α_{ml} and the RHD T(τ) relations provides the same $\alpha_{ml,\odot}=1.82$.

Solar calibrations with the KS, VAL_c and Eddington T(τ) relations have provided $\alpha_{ml,\odot}$ =2.11, 1.90 and 1.69, respectively.

In all these calibrated solar models the initial solar He mass fraction ($Y_{\text{ini},\odot} \sim 0.274$) and metallicity (($Z_{\text{ini},\odot} \sim 0.0199$) are essentially the same, as expected. Also, the model present He mass fraction in the envelope ($Y_{\odot} = 0.244$) and the depth of the convection zone ($d_{CZ} = 0.286R_{\odot}$) are the same for all calibrations and in agreement with the helioseismic values (see, e.g., Dziembowski et al. 1995; Basu & Antia 1997).

If we take $\alpha_{ml,\odot}$ obtained with the RHD T(τ) as a reference, the lower value obtained with the Eddington T(τ), and the larger values obtained with both the VAL_c and KS relations (in increasing order) are fully consistent with the results of Fig. 3. In that figure VAL_c and KS T(τ) MS models are increasingly hotter than the reference RHD calculations (hence increasingly larger α_{ml} values are required to match the reference MS) whereas the use of the Eddington T(τ) produces models cooler than the reference MS (hence lower α_{ml} value are needed to match the reference MS).

4. Summary and discussion

We have presented the first self-consistent stellar evolution calculations that employ the variable α_{ml} and $T(\tau)$ by Trampedach et al. (2014b), based on their 3D RHD simulations. Our set of evolutionary tracks for different masses and the same chemical composition (plus consistent Rosseland opacities) of the RHD simulations, cover approximately the entire g- T_{eff} parameter space of the 3D atmosphere/envelope calculations.

We found that, from the point of view of the predicted $T_{\rm eff}$ (plus the depth of the convective envelopes and amount of pre-MS Li depletion), models calculated with constant RHD calibrated $\alpha_{\rm ml,\odot} = 1.76$ are very close to, and often indistinguishable from, the models with variable $\alpha_{\rm ml}$. Maximum differences are at most ~30-50 K. This result is similar to the conclusions by Freytag & Salaris (1999), based on 2D RHD simulations at low metallicities.

At first sight this may appear surprising, given that the full range of α_{ml} spanned by the RHD calibration is between ~1.6 and ~2.0. However, one has to take into account the following points:

- 1. The derivative $\Delta T_{\rm eff}/\Delta \alpha_{\rm ml}$ for stellar evolution models depends on the absolute value of α_{ml} , and decreases when α_{ml} increases).
- 2. The variation of $\Delta T_{\rm eff}$ for a given $\Delta \alpha_{\rm ml}$ depends also on the mass extension of the convective and superadiabatic regions.

It is therefore clear that the decrease of $\alpha_{\rm ml}$ with increasing $T_{\rm eff}$ cannot have a major effect because of the thin convective (and superadiabatic) layers of models crossing this region of the g- $T_{\rm eff}$ diagram. The large variations $\Delta \alpha_{\rm ml} \sim 0.2$ (see lower panel of Fig. 1) along the lower Hayashi track of the $1.0M_{\odot}$ model is also not very significant (~50 K) because of the decreased extension of the surface convection and the reduced $\Delta T_{\rm eff}/\Delta \alpha_{\rm ml}$ at higher α_{ml} .

It is however very important to remark here that the detailed structure of the superadiabatic convective regions is not suitably reproduced either by $\alpha_{ml,\odot}$ or by a variable α_{ml} , and that the full results from RHD models need to be employed whenever a detailed description of the properties of these layers is needed.

To some degree the role played by the RHD $T(\tau)$ is more significant. Constant $\alpha_{ml,\odot}$ = 1.76 stellar evolution models become systematically cooler by up to ~100 K along the MS, pre-MS and RGB when the widely used KS $T(\tau)$ relation is used, compared to the self consistent RHD-calibrated calculations. Evolutionary tracks obtained employing the Eddington and VAL_c $T(\tau)$ relationships are much less discrepant, and stay within ~50 K of the self-consistent calculations. Even from the point of view of pre-MS Li depletion the VAL_c $T(\tau)$ causes only minor differences, of the order of 10%. A similar difference is found with the KS relation, whilst a larger effect of ~20% is found with the Eddington $T(\tau)$.

An extension of Trampedach et al. (2013) simulations to different metallicities is necessary to extend this study and test the significance of the α_{ml} variability (and Hopf functions) over a larger parameter space. The 3D RHD simulations by Magic et al. (2013) and the α_{ml} calibration by Magic et al. (2015) cover a large metallicity range, and predict an increase of α_{ml} with decreasing metal content, but at the moment it is not possible to properly include this calibration in stellar evolution calculations, for the lack of available $T(\tau)$ relations and input physics consistent with the RHD simulations. At solar metallicity the general behaviour of α_{ml} with g and T_{eff} seems to be qualitatively similar to Trampedach et al. (2014b) calibration. However, the range of α_{ml} is shifted to higher values compared to Trampedach et al. (2014b) results, due probably to different input physics and the different adopted solar chemical composition.

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