Equation of state for solar near-surface convection

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Abstract. Numerical 3-D radiative hydrodynamical simulations are the main tool for the analysis of the interface between the solar convection zone and the photosphere. The equation of state is one of the necessary ingredients of these simulations. We compare two equations of state that are commonly used, one ideal and one nonideal, and quantify their differences. Using a numerical code we explore how these differences propagate with time in a 2-D convection simulation. We show that the runs with different equations of state (EOSs) and everything else identical relax to statistically steady states in which the mean temperature (in the range of the continuum optical depths typical for the solar photosphere) differs by less than $0.2\%$. For most applications this difference may be considered insignificant.

Keywords. Solar physics astrophysics and astronomy (photosphere and chromosphere)

1 Introduction

Realistic time-dependent 3-D numerical simulations (Nordlund, 1982; Stein and Nordlund, 1998, 2000) radically improved our knowledge of the near-surface convection and the solar photosphere. In these simulations the equations of magnetohydrodynamics, the equation of state (EOS) and the radiative transfer equation (RTE) are solved ab initio with only few free parameters. Apart from the three-dimensionality, the key ingredient that makes these simulations realistic is the detailed description of the microphysical processes contributing to the EOS and the RTE. The realism of the simulations had been repeatedly confirmed by diverse numerical experiments comparing the actual observations of the sun with the spectra computed a posteriori from the models (Asplund et al., 2000b; Khomenko et al., 2005; Shelyag et al., 2007; Danilovic et al., 2008; Wedemeyer-Böhm and Rouppe van der Voort, 2009; Pereira et al., 2013).

Beeck et al. (2012) compared the properties of the 3-D hydrodynamical (HD) simulations carried out by three codes that are the workhorses in the field: Stagger (Galsgaard and Nordlund, 1996), MPS/University of Chicago Radiative Magnetohydrodynamics (MURaM) (Vögler et al., 2005) and COnservative COde for the COmputation of COmpressible COnvection in a BOx of L Dimensions, $L = 2, 3$ (Co5Bold) (Wedemeyer et al., 2004). The codes are all primarily designed for simulations of near-surface convection, but with some intrinsic differences in the details of the microphysics, in the numerical algorithms, the boundary conditions, the parallelisation strategy, etc. Nevertheless, Beeck et al. (2012) demonstrated that the mean properties and the spatial distribution of the fundamental quantities in the HD simulations performed with these codes do not vary significantly. The difference in the results of these simulations is attributed to the difference in the input parameters (the opacities, the chemical composition). However, because the outcome of the code depends on these parameters in a highly non-linear way, it is impracticable to trace back the differences to the initial assumptions and approximations. An independent code was developed more recently by Tanner et al. (2012, the RHD code). Although the RTE in this code is solved in a simplified way, the authors showed that their results of the convection simulation are comparable to the results of Beeck et al. (2012). Tanner et al. (2013) employed the radiative hydrodynamics (RHD) code to study the variation in the simulated stellar convection with varying input metallicity. They concluded that the variation in the superadiabaticity and the convection dynamics in their simulation remains relatively small for a large range of metallicities ($0.01 \leq Z \leq 0.40$).
The case of the pure hydrogen plasma (after Hansen et al., 2004). The boundaries are estimated for the partial ionisation effects and (2) the ideal EOS for a mixture including the pressure ionisation, Coulomb interaction and electron degeneracy. MHD degeneracy and (2) the ideal EOS for a mixture including the pressure ionisation, Coulomb interaction and electron degeneracy.

Two types of EOS are used in the codes for the solar convection simulations: (1) the nonideal EOS including the effects of the pressure ionisation, Coulomb interaction and electron degeneracy and (2) the ideal EOS for a mixture including the partial ionisation effects. MHD$^1$ (Hummer and Mihalas, 1988; Mihalas et al., 1988) and OPAL (Rogers et al., 1996; Rogers and Nayfonov, 2002) are commonly used nonideal EOSs. The former is used in Stagger, the latter in MURaM, RHD, in the ANTARES code (Grimm-Strele et al., 2015) and in the MANCHA code (Khomenko et al., 2014). The ideal EOS of Wolf (1983) is implemented in Co5Bold. MURaM can work with a simple ideal EOS described in Vögler et al. (2005), while MANCHA incorporates an algorithm for the evaluation of the ideal EOS of a partially ionised mixture described by Vardya (1965) and improved by Mihalas (1967) and Wittmann (1974) (we shall refer to this EOS as VMW). The VMW algorithm is implemented in several radiative transfer codes to provide the electron pressure for known temperature and gas pressure (e.g. Bellot Rubio et al., 1999; Socas-Navarro, 2011).

The range of validity of the ideal EOS may be estimated for the trivial case of partially ionised pure hydrogen. In Fig. 1, we reproduce Fig. 3.9 of Hansen et al. (2004), showing the $T$–$\rho$ plane with the boundaries for the nonideal effects and the curve where the ionisation fraction of hydrogen is 0.99. To Fig. 1 we add plots of two 1-D stratifications (the chromospheric VALC model of Vernazza et al. (1981) and the convection zone model of Spruit, 1974) and the area covered by a typical 3-D HD simulation (shaded area). All three models are within the limits of where the ideal EOS for pure hydrogen is valid. The deep end of the convection-zone model is close to the pressure–ionisation boundary, while, on the other end, the ionisation of hydrogen in the chromosphere requires a non-equilibrium solution of hydrogen ionisation (Carlsson and Stein, 2002) that cannot be represented by a simple curve in this plot. Nevertheless, the near-surface convection and the photosphere fall in the region of the $T$–$\rho$ plane, where the ideal EOS is a safe assumption.

To compute nonideal EOS is extremely time-consuming and impossible to do on the fly in codes for 3-D numerical simulations. Instead EOS is precomputed and results are stored in lookup tables suitable for fast interpolation. The computational cost of the ideal EOS is much lower; however, when the molecules are taken into account, it is necessary to solve the equations iteratively, and, therefore, the use of the precompiled lookup tables may save considerable computing time as long as the table grid is sufficiently fine to limit the interpolation errors.

2 Ideal versus nonideal EOS

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2.1 Ideal EOS for partially ionised mixture

The VMW EOS accounts for the partial ionisation of the mixture composed of atomic and molecular hydrogen (H, H$^-$, H$^+$, H$_2$ and H$_2^+$), neutral, singly and doubly ionised atoms of helium and other elements. The ionisation equilibrium is computed using Saha’s equation, while for the dissociation of the molecules, the instantaneous chemical equilibrium is assumed. We implemented the algorithm in a stand-alone code with several improvements, mainly related to the atomic data. For the atomic partition functions, we adopted polynomial fits of Irwin (1981). The molecular
The ionisation fraction of hydrogen in the $T$–$p$ grid computed with the VMW EOS. Those portions of the domain that are dominated by the neutral H, H$_2$ molecule and H$^+$ are indicated.

Figure 2. The ionisation fraction of hydrogen in the $T$–$p$ grid computed with the VMW EOS. Those portions of the domain that are dominated by the neutral H, H$_2$ molecule and H$^+$ are indicated.

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2.2 Nonideal EOS: OPAL

The OPAL EOS is computed in the physical picture through an activity expansion of the grand canonical partition function of the plasma. The results are distributed as lookup tables of the internal energy, the pressure, the electron density and other derived thermodynamical quantities as functions of the temperature and the density of several chemical compositions. We reversely interpolate the tables (for the mass fraction $X = 0.7062$ and $Z = 0.0197$, corresponding to the abundances of Anders and Grevesse) to obtain the internal energy, the density and the electron pressure in the $T$–$p$ coordinates. The interpolation is performed using the method of the overlapping biquadratics that is included in the distribution of the OPAL EOS. The results for the internal energy and the density shown in the same range of the temperature and pressure as in Fig. 3 are visually nearly identical to the results of VMW. The electron density distribution shows a number of artefacts in the low-temperature region where ad hoc electron density is introduced (Rogers and Nayfonov, 2002), while it is similar to the VMW distribution elsewhere. Trampedach et al. (2006) examined the differences between the OPAL EOS and the MHD EOS.

2.3 Comparison

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The relative differences between the mass density, the electron number density and the internal energy in the VMW and the OPAL EOS in the $\log T$–$\log p$ plane are shown in Fig. 4. The positive values indicate that the VMW quantity is larger. The largest differences in the density and the internal energy occur in the area with a temperature below $10^6$ K for pressure higher than $10^9$ dyn cm$^{-2}$ and densities higher than $10^{-6}$ g cm$^{-3}$. This is the area where hydrogen is still neutral in the VMW EOS (cf. Fig. 2). The OPAL tables do not include the partial pressures of different particles needed for a direct comparison. However, the negative difference (the red area) in the density at constant pressure together with correspondingly higher internal energy and the electron density indicate that the VMW hydrogen ionisation fraction is higher than the one from OPAL. The opposite applies for the blue area. While the latter may be interpreted as the effect of the pressure ionisation (cf. Fig. 1), it is difficult to attribute the former to any particular effect without knowing the partial pressures consistent with the OPAL. Nevertheless, the area where the differences appear is not covered by the near-surface convection. At low pressure and a temperature below $\approx 2 \times 10^4$ K, the relative differences are below 1% (except for the electron densities at low $T$, where the artefacts dominate). Figure 5 shows a blow-up of Fig. 4 in that region. Vertical stripes in the density plot and the horizontal stripes in the electron pressure are due to the interpolation errors (coming from the reverse interpolation of the OPAL table to the $T$–$p$ domain)$^4$. The vertical bands below 3500 K in the relative difference of the electron density correspond

$^2$Hereafter we use “internal energy” for the specific internal energy per mass.

$^3$http://opalopacity.llnl.gov/EOS_2005/

$^4$Note that we use overlapping biquadratics for the interpolation as in the code included in the OPAL distribution. The interpolation errors may be suppressed by a different interpolation scheme, e.g. with bicubic splines.
to the area where the OPAL electron densities are unreliable.

3 The convection simulation

To study propagation with time of differences caused by EOS choice in a near-surface convection simulation, we initiate and run two simulations with all parameters identical except for the EOS lookup tables. For this experiment we use the version of the MURaM code described in Vögler (2003) with several minor adaptations needed to run the code with an alternative set of EOS lookup tables. The code requires two tables, one with the temperature and the density tabulated as functions of the internal energy and the pressure and another with the temperature and the pressure as functions of the internal energy and the density. The required tables are produced from the VMW and the OPAL EOS described above. Since we are primarily interested in the relative difference between the two runs and not in the physical realism of the result, we set up a simple HD convection run in 2-D with grey opacities. The physical size of the domain is 6.0 Mm in the horizontal direction and 1.4 Mm in the vertical, with 288 and 100 grid points respectively. The bottom boundary is open for the mass flow, the top boundary is closed and the vertical boundaries are set as periodic. The mass fluctuations through the bottom boundary are dynamically adjusted by the boundary pressure of the upflowing material \( p_{up} \) (see Vögler, 2003, Eq. 3.56, p. 32).

The initial model is prepared by replicating the temperature and the density of a 1-D convectively stable model in two dimensions. The magnetic field is equal to 0 throughout the simulation runs. The internal energy is computed from these two quantities using both VMW and OPAL, so that we have two initial snapshots, one consistent with each EOS.
Figure 5. Blow-up of Fig. 4 in the low-pressure and low-temperature region relevant for the near-surface convection. The fragments of the contour lines from Fig. 4 are added.

Figure 6. Vertical velocity at fixed height (850 km above the bottom of the simulation box) over the first 24 min of the simulation with the OPAL EOS (left) and with the VMW EOS (right). The dashed line approximately marks the instance when the two simulation runs separate.

The specific energy in the snapshots is then perturbed with a multiplicative factor of $1 \pm 0.05$, with identical randomisation over the simulation box. We start the runs with a damping factor for the radiative exchange term, gradually releasing it until the convection cells fully develop. Once the simulation is in the statistically steady state, we let it run for about 16 h of solar time while taking a snapshot every 20 s.

3.1 Results

In the first iteration step, the relative difference between the temperature in the two runs is within $\pm 0.2\%$, which is in agreement with Fig. 5. As the flows develop and accelerate in the initial phase, the differences between the runs grow and the point-by-point comparison between their parameters becomes meaningless. Figure 6 shows the velocity for the two runs at constant height in the box (850 km above the bottom of the box) versus time. The flows in the two cases are indistinguishably similar over the first 13 min; after that initial phase they take different evolutionary paths toward the statistically steady solutions.

In MURaM the flux of the emergent radiation $F_{\text{out}}$ is controlled by the specific energy of the upflowing material at the bottom boundary so that its mean value matches the value of the real sun (see Vögler, 2003, Eq. 3.50, p. 30). The relative difference of the mean $F_{\text{out}}$ for the two runs is thus small, as expected: 0.3 %. The temperature profiles of the two runs averaged over time and over the horizontal direction are very similar. The difference in the geometrical height scale (left panel) and in the scale of the continuum optical depth at 500 nm (right) is shown in Fig. 7. The largest absolute difference between the two in the geometrical height scale is at 0.8 Mm above the bottom of the box and amounts to nearly 200 K (the relative difference is 2.3 %). This is the region where the radiative cooling begins to dominate producing a steep gradient of temperature with height. In the continuum optical depth scale at 500 nm, in the photosphere ($\tau_{500} \leq 1$), the difference between the two temperature profiles is up to 10 K (less than 0.2 % relative to the local temperature). This difference causes the 0.3 % difference in the mean flux of the emergent intensity, which is negligible for most purposes.
4 Conclusions

The tables of thermodynamical quantities (the mass density, the specific internal energy and the electron density) computed using two equations of state – one ideal (VMW, after Vardya, 1965; Mihalas, 1967; Wittmann, 1974) and one nonideal (OPAL, Rogers et al., 1996; Rogers and Nayfonov, 2002) and both widely used in solar physics – are compared. The relative differences between the quantities in the region of the $T$–$p$ plane usually considered in the realistic near-surface convection simulations (approximately 1 Mm below the level where the mean continuum optical depth is equal to 1 and 0.5 Mm above it) are below 1%, except for the electron density at the low temperatures, where the values of the OPAL tables are not reliable. Since we have access only to the tables precomputed with the OPAL code but not to the code itself, we cannot determine the origin of these differences. Possible causes are the differences in the atomic and molecular data, in the approach and the interpolation errors.

To check how this difference may affect the result of the hydrodynamical simulations, we simulated the 2-D solar convection with the two EOSs with an otherwise identical set-up. The two simulation runs separate in the parameter space as soon as the convective cells are developed. Nevertheless, they evolve separately to a statistically nearly identical steady state, with the temperature difference in the optical depth scale below 10 K throughout the photosphere. This experiment demonstrates that the two EOSs are interchangeable for simulations of near-surface solar convection. Moreover, the electron density in the OPAL tables, corrupted at low temperatures, may be recomputed for a large portion of the $T$–$\rho$ grid of OPAL with an accuracy better than 1% using the VMW ideal EOS (cf. Fig. 4). To estimate what the effect of the EOS choice in a more realistic set-up is (3-D, with magnetic field, non-grey radiative transfer including scattering, etc.), it would be necessary to repeat this experiment in such conditions. The change of the mean temperature between 2-D and 3-D solar convection simulations has been described by Asplund et al. (2000a) (see lower panel of their Fig. 9). Without analysing the origin of the differences in their study, it may be noted that these differences are of the same order as the differences between the highly realistic 3-D HD simulations performed by different codes (Beeck et al., 2012). In any case, we do not expect significant changes in our results as long as the temperature and the density in a simulation remain within nearly the same range of the $T$–$\rho$ plane as in our experiment.

This result is in agreement with the studies of Beeck et al. (2012) and Tanner et al. (2012), demonstrating robustness of the near-surface simulations. Regarding the deep convection, this result is consistent with the conclusions of Bahcall et al. (2004) that the depth of the convection zone does not depend significantly on the uncertainties in the EOS. It is also consistent with the recent study of Lord et al. (2014), who compared the effect of the OPAL EOS and a simple Saha-based EOS on the horizontal velocity spectrum of a solar deep convection simulation and found that the two produced nearly identical results, especially in the higher portion of the convection zone.

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