Radiation-driven winds of hot luminous stars
XIII. A description of NLTE line blocking and blanketing
towards realistic models for expanding atmospheres

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Abstract. Spectral analysis of hot luminous stars requires adequate model atmospheres which take into account the effects of NLTE and radiation driven winds properly. Here we present significant improvements of our approach in constructing detailed atmospheric models and synthetic spectra for hot luminous stars. Moreover, as we regard our solution method in its present stage already as a standard procedure, we make our program package WM-basic available to the community (download is possible from the URL given below). The most important model improvements towards a realistic description of stationary wind models concern:

(i) A sophisticated and consistent description of line blocking and blanketing. Our solution concept to this problem renders the line blocking influence on the ionizing fluxes emerging from the atmospheres of hot stars – mainly the spectral ranges of the EUV and the UV are affected – in identical quality as the synthetic high resolution spectra representing the observable region. In addition, the line blanketing effect is properly accounted for in the energy balance.

(ii) The atomic data archive which has been improved and enhanced considerably, providing the basis for a detailed multilevel NLTE treatment of the metal ions (from C to Zn) and an adequate representation of line blocking and the radiative line acceleration.

(iii) A revised inclusion of EUV and X-ray radiation produced by cooling zones which originate from the simulation of shock heated matter.

This new tool not only provides an easy-to-use method for O-star diagnostics, whereby physical constraints on the properties of stellar winds, stellar parameters, and abundances can be obtained via a comparison of observed and synthetic spectra, but also allows the astrophysically important information about the ionizing fluxes of hot stars to be determined automatically. Results illustrating this are discussed by means of a basic model grid calculated for O-stars with solar metallicity. To further demonstrate the astrophysical potential of our new method, we first provide a detailed spectral diagnostic determination of the stellar parameters, the wind parameters, and the abundances by an exemplary application to one of our grid-stars, the O9.5Ia O-supergiant α Cam. Our abundance determinations of the light elements indicate that these deviate considerably from the solar values.

Key words. line: formation – stars: atmospheres – stars: early type – stars: mass-loss – stars: individual: α Cam – X-rays: stars

1. Introduction

Spectral analyses of hot luminous stars are of growing astrophysical interest as they provide a unique tool for the determination of the properties of young populations in galaxies. This objective, however, requires spectral observation of individual objects in distant galaxies. That this is feasible has already been shown by Steidel et al. (1996) who detected galaxies at high redshifts (z ~ 3.5) and found that the corresponding optical spectra show the typical features usually found in the UV spectra of hot stars. In order to determine stellar abundances and physical properties of the most UV-luminous stars in at least the Local Group galaxies via quantitative UV spectroscopy, another principal difficulty needs to be overcome: the diagnostic tools and techniques must be provided. This requires the construction of detailed atmospheric models and synthetic spectra for hot luminous stars. It is a continuing effort of several groups to develop a standard code for solving this problem. Recent basic papers of the different groups concerning O-stars are Pistinner et al. (1999), Aufdenberg et al. (1998), Schaerer & de Koter (1997), Pauldrach et al. (1994a, 1994b, 1998),

The most important output of this kind of model calculation are the ionizing fluxes and synthetic spectra emitted by the atmospheres of hot stars. As these spectra consist of hundreds of not only strong, but also weak wind-contaminated spectral lines which form the basis of a quantitative analysis, and as the energy distribution from hot stars is also used as input for the analysis of emission line spectra (e.g., of gaseous nebulae) which depend sensitively on the structure of the emergent stellar flux, a sophisticated and well tested method is required to produce these data sets accurately.

However, developing such a method is not straightforward, since modelling hot star atmospheres involves replicating a tightly interwoven mesh of physical processes: the equations of radiation hydrodynamics including the energy equation, the rate equations for all important ions (from H to Zn) including the atomic physics, and the radiative transfer equation at all transition frequencies have to be solved simultaneously.

The most complicating effect in this system is the overlap of thousands of spectral lines of different ions. Especially concerning this latter point we have made significant progress in developing a fast numerical method which accounts for the blocking and blanketing influence of all metal lines in the entire sub- and supersonically expanding atmosphere.

As we have found from previous model calculations that the behavior of most of the UV spectral lines depends critically on a detailed and consistent description of line blocking and line blanketing (cf. Pauldrach 1987; Pauldrach et al. 1990; Pauldrach et al. 1994a; Sellmaier et al. 1996; Taresch et al. 1997; Haser et al. 1998; this has also been pointed out by Schaerer & Schmutz 1994; Schaerer & de Koter 1997; Hillier & Miller 1998), special emphasis has been given to the correct treatment of the Doppler-shifted line radiation transport, the corresponding coupling with the radiative rates in the rate equations, and the energy conservation.

In Sect. 3 we will demonstrate that the realistic and consistent description of line blocking and blanketing and the modifications to the models involved lead to changes in the energy distributions, ionizing continua, and line spectra with much better agreement with the observed spectra when compared to previous, not completely consistent models. This will obviously have important repercussions for the quantitative analysis of hot star spectra.

In the next two sections we will first summarize the general concept of our procedure and then discuss the current status of our treatment of hydrodynamical expanding atmospheres.

2. The general method

The basis of our approach in constructing detailed atmospheric models for hot luminous stars is the concept of homogeneous, stationary, and spherically symmetric radiation driven winds, where the expansion of the atmosphere is due to scattering and absorption of Doppler-shifted metal lines (Lucy & Solomon 1970). In contrast to previous papers of this series, the above approximations are now the most significant ones for the present approach. These approximations are, however, quite restrictive, since only the time-averaged mean of the observed spectral features can be described correctly by our method. Nevertheless we believe that it is reasonable to continue with the stationary, spherically symmetric approach and to improve its inherent physics, since the detailed comparison with the observations, which is the only way to demonstrate the reliability of this concept, leads to promising results (cf. Sect. 4).

Before we describe the latest improvements in detail we first summarize the principal features of our procedure of simulating the atmospheres of hot stars. (For particular points, a comprehensive discussion is also found in the papers cited above.)

Figure 1 gives an overview of the physics to be treated in various iteration cycles. A complete model atmosphere calculation consists of three main blocks,

(i) the solution of the hydrodynamics;
(ii) the solution of the NLTE-model (calculation of the radiation field and the occupation numbers);
(iii) the computation of the synthetic spectrum.

which interact with each other.

In the first step the hydrodynamics is solved in dependence of the stellar parameters (effective temperature \( T_{\text{eff}} \), surface gravity \( \log g \), stellar radius \( R_\star \) (defined at a Rosseland optical depth of 2/3), and abundances \( Z \) (in units of the corresponding solar values) and of pre-specified force multiplier parameters \( (\delta_0, \alpha_0, \delta_0) \), which are used for describing the radiative line acceleration. In addition, the continuum force is approximated by the Thomson force, and a constant temperature structure \( T(r) = T_{\text{eff}} \) is assumed in this step. In a second step the hydrodynamics is solved by iterating the complete continuum force \( g_C(r) \) (which includes the opacities of all important ions) and the temperature structure (both are calculated using a spherical grey model), and the density \( \rho(r) \) and the velocity structure \( v(r) \). In a final outer iteration cycle these structures are again iterated together with the line force \( g_L(r) \) obtained from the spherical NLTE model. (New force multiplier parameters \( (k, \alpha, \delta) \), which are depth dependent if required, are deduced from this calculation\(^1\)).

The main part of the code consists of the solution of the NLTE-model. In this step the radiation field (represented by the Eddington-flux \( H_{\nu}(r) \) and the mean intensity \( J_{\nu}(r) \), the final temperature structure \( T(r) \), occupation numbers \( n_i(r) \), and opacities \( \chi_\nu \) and emissivities \( \eta_\nu \) are

\(^1\) This latter step is currently not available for the download version of the code; it will be made available for version 2.0.
Hydrodynamics

- Line force: $g_L(k, \alpha, \delta)$
- Continuum force: $g_C(r)$
- Temperature: $T(r)$

\[ \rho(r), v(r) \implies \dot{M}, v_\infty \]

Spherical grey model

- $\rho(r), v(r)$
- LTE continuum opacities

\[ T_g(r), g_C(r) \implies \text{LTE continuum force}
\text{LTE temperature} \]

Spherical NLTE model

\[ Z, \rho(r), v(r), T_g(r), R_{ij}, C_{ij}, \chi_\nu, \eta_\nu \]

\[ n_i(r), H_\nu(r), T(r), g_L(r), g_C(r) \implies \text{line force}
\text{continuum force}
\text{temperature} \]

Spectrum (formal solution)

\[ n_i(r), \chi_\nu, \eta_\nu, v(r) \]

\[ H_\nu^m \implies \text{line and continuum}
\text{spectrum} \]

Fig. 1. Sketch of a complete model atmosphere calculation. Starting procedures are presented in brackets. For a discussion see the text.

Computed using detailed atomic models for all important ions. For the solution of the radiative transfer equation, the influence of the spectral lines (i.e., the UV and EUV line blocking) is properly taken into account in addition to the usual consideration of continuum opacities and source functions consisting of Thomson-scattering and free-free and bound-free contributions of all important ions. Moreover, the shock source functions produced by radiative cooling zones which originate from a revised simulation of shock heated matter are also included. For the calculation of the final NLTE temperature structure the line blanketing effect, which is a direct consequence of line blocking, is considered by demanding luminosity conservation and the balance of microscopic heating and cooling rates. The rate equations which yield the occupation numbers contain collisional ($C_{ij}$) and radiative ($R_{ij}$) transition rates, as well as low-temperature dielectronic recombination and Auger-ionization due to K-shell absorption (considered for C, N, O, Ne, Mg, Si, and S) of soft X-ray radiation arising from shock-heated matter. (Further details concerning the solution method of the NLTE-model are described in Sect. 3.)

The last step consists of the computation of the synthetic spectrum for the purpose of comparison with observations. In dependence of the occupation numbers, the opacities and the emissivities, a formal integral solution of
the transfer equation in the observer’s frame is performed (cf. Puls & Pauldrach 1990).

As results of the iterative solution of this system of equations we obtain not only the synthetic spectra and ionizing fluxes which can be used in order to determine stellar parameters and abundances, but also the hydrodynamical structure of the wind (thus, constraints for the mass-loss rate \( \dot{M} \) and the velocity structure \( v(r) \) can be derived).

3. The consistent NLTE model

The construction of realistic models for expanding atmospheres requires a correct and completely consistent description of the main part of the simulation, the NLTE model. In this regard, the most crucial point in our present improved treatment is an exact description of line blocking and blanketing.

The effect of line blocking – mainly acting in between the He II and the H I edge – is that it influences the ionization and excitation and the momentum transfer of the radiation field significantly. This of course has important consequences for both the spectral line formation and the dynamics of the expanding atmosphere. Nevertheless, it is still not a common procedure to treat the line opacities and emissivities in the radiative transfer equation and their back-reaction on the occupation numbers via the radiative rates correctly. We will therefore first discuss the effects of line blocking and blanketing for expanding atmospheres of hot stars in more detail.

The huge number of metal lines present in hot stars in the EUV and UV attenuate the radiation in these frequency ranges drastically by radiative absorption and scattering processes (an effect known as line blocking). Only a small fraction of the radiation is re-emitted and scattered in the outward direction; most of the energy is radiated back to the surface of the star producing there a backwarming. Due to the increase of the Rosseland optical depth \( \tau_{\text{Ross}} \) resulting from the opacities enhanced by the line blocking, and, in consequence, of the temperature, the radiation is redistributed to lower energies (this refers to line blanketing). In principle these effects influence the NLTE model with respect to:

(i) the radiative photoionization rates \( R_{i \alpha} \);
(ii) the radiative bound-bound rates \( R_{ij} \);
(iii) the radiation pressure \( g_{\text{rad}} \);
(iv) the energy balance.

The terms of the first two items are directly connected to the radiation field, and line blocking in general reduces them considerably. Concerning the third item, the blocked incident radiation reduces the radiative acceleration term in the inner part, whereas it can be enhanced in the outer part due to multiple scattering processes (cf. Puls 1987 and references therein). In contrast to this, the energy equation – last item – is mostly influenced by the impact of the line opacities, and this blanketing effect results in an increased temperature (steeper gradient) in the deeper layers of the photosphere.

Although the method for treating blanketing effects is well established for cold stars, where the atmospheres are hydrostatic and where the assumption of LTE is justified (cf. Kurucz 1979 and 1992), the work to develop an adequate method for hot stars, where not only NLTE effects are prominent, but where the atmospheres are also rapidly expanding, is still under way. (For the various approaches taken to this end, see the references listed in Sect. 1.) In this case – hot stars with expanding atmospheres – in addition to the four items given above, the solution of the radiative transfer also has to account for the lineshift caused by the Doppler effect due to the velocity field. The important effect of this point is that the velocity field increases the frequential range which can be blocked by a single line (see below). In the presence of a velocity field the blocking effect is therefore more pronounced.

Concerning the basic requirements for calculating adequate line opacities and source functions for expanding atmospheres of hot stars we have to concentrate on the following points:

(1) consistent NLTE occupation numbers;
(2) a complete and accurate line list in connection with detailed atomic models;
(3) a proper concept for treating the line blocking with due regard to the lineshifts in the wind, in the course of which the method for solving the complete radiative transfer including the spectral lines has to be efficient with regard to computational time;
(4) a correct treatment of the influence of the blanketing effect on the temperature structure;
(5) an adequate approximation of the EUV and X-ray radiation produced by cooling zones of shock-heated matter.

3.1. The concept of the solution of ionization and excitation

It is obvious that ionization and excitation plays the major role in calculating the emergent flux and spectrum of a hot star. Therefore, a consistent and accurate description of the occupation numbers is extremely important for a realistic solution of the NLTE model.

Figure 2 presents a sketch of our iteration scheme for the calculation of the occupation numbers. To save more than a factor of 20 in computation time, the iteration is performed in two major steps, which differ mainly in the accuracy achieved by the methods employed: in a pre-iteration, a modified opacity sampling technique (method I) is used to take into account in the solution of the radiation transfer the hundreds of thousands of spectral lines in the UV and EUV. The main requirements for this step are that it is sufficiently accurate for the iteration to converge near the final solution, but fast enough to make the model calculation feasible with today’s computers. In the final iterations, the radiation transfer (taking into account the same lines as in method I) is calculated...
with an exact solution of the transfer equation in the observer’s frame (method II). The advantage of this method is that it is free from all major approximations; its disadvantage is its comparatively high computational cost. As the pre-iteration has already converged near the final solution, however, only very few of these final iteration steps are needed. Details are discussed in the following paragraphs.

Note that both methods are used successively – method I for at least 500–600 iterations, and then method II for at least another 30–150 iterations – and the same quantities are iterated (see Fig. 2). Both methods are of course based on the same radius and frequency grids and take into account the same lines. The whole purpose of method I is to give good starting values for the final (real) iteration cycle using method II. In fact, as will be shown below, the starting values produced by method I turned out to be rather excellent.

In dependence of the abundances \((Z)\), the density \((\rho(r))\) and velocity \((v(r))\), and a pre-specified temperature structure \((T_g(r))\) (see Sect. 2), the occupation numbers are determined by the rate equations containing collisional \((C_{ij})\) and radiative \((R_{ij})\) transition rates. The most crucial dependency of the rates is not the density, which is nevertheless important for the collisional rates and the equation of particle conservation, but the velocity field which enters not only directly into the radiative rates via the Doppler shift, but also indirectly through the radiation field determined by the equation of transfer, which in turn is again dependent on the Doppler shifted line opacities and emissivities.

For the calculation of the radiative bound-bound transition probabilities \(R_{ij}\) we make use of the Sobolev plus-continuum method (Hummer & Rybicki 1985; Puls & Hummer 1998). Only for some weak second-order lines in the subsonic region of the atmospheric layers where the continuum is formed might this be just a poor approximation (cf. Sellmaier et al. 1993). A more important point of our procedure concerns the problem of self-shadowing (cf. Pauldrach et al. 1998). This problem
occurs because the rate equations are not really solved simultaneously with the radiative transfer, but instead in the framework of the accelerated lambda iteration (ALI), in which the radiation field and the occupation numbers are alternately computed (cf. Pauldrach & Herrero 1988). Hence, the radiation field which enters into a bound-bound transition probability is already affected by the line itself, since the line has also been considered for the blocking opacities. This procedure will lead to a systematic error if a line transition dominates within a frequency interval (see Sect. 3.3.1). The solution for correctly calculating the bound-bound rates even in these circumstances is quite simple and has been described by Pauldrach et al. (1998, Sect. 3.2).

The spherical transfer equation yields the radiation field at 2500 frequency points (see below) and at every depth point, including the layers where the radiation is thermalized and hence the diffusion approximation is a proper boundary condition. The solution includes all relevant opacities. In particular, the effects of wind and photospheric EUV line blocking on the ionization and excitation of ions of the 149 ionization stages of the 26 elements (both continuum and lines) and the resulting occupation numbers and of the synthetic spectrum is directly dependent on the quality of the input data. We have therefore extensively revised and improved the basis of our model calculations, the atomic models.

Up to now, the atomic models of all of the important ions of the 149 ionization stages of the 26 elements considered (H to Zn, apart from Li, Be, B, and Sc) have been replaced in order to improve the quality. This has been done using the Superstructure program (Eissner et al. 1974; Nussbaumer & Storey 1978), which employs the configuration-interaction approximation to determine wave functions and radiative data. The improvements include more energy levels (comprising a total of about 5000 observed levels, where the fine structure levels have been “packed” together2) and transitions (comprising more than 30000 bound-bound transitions for the NLTE calculations and more than 400000 lines for the line-force and blocking calculations3,4), and 20000 individual transition probabilities of low-temperature dielectronic recombination and autoionization.

Regarding the latter point, the inclusion of line opacities and emissivities in the transfer equation, two different concepts are employed for iterating the occupation numbers and the temperature structure until a converged radiation field \((J_\nu(r) \text{ and } H_\nu(r))\) is obtained. In a first step, a pre-iteration cycle with an opacity sampling method is used (method I). This procedure has the advantage of only moderate computing time requirements, allowing us to perform the major part of the necessary iterations with this method. Its disadvantage, however, is that it involves a few substantial approximations (cf. Sect. 3.3). In a second step, the final iteration cycle is therefore solved with the detailed radiative line transfer (method II). Although this procedure is extremely time-consuming, it has the advantage that it is not affected by any significant approximations. With this second method, blocking factors \(B_J(r, \nu)\) and \(B_H(r, \nu)\) are calculated, defined as the ratio of the radiative quantities obtained by considering the total opacities and emissivities to those which include only the corresponding continuum values (cf. Pauldrach et al. 1996). \(B_J(r, \nu)\) and \(B_H(r, \nu)\) are then used as multiplying factors to the continuum quantities calculated in the next NLTE-ALI-cycle with the current continuum opacities, in order to iterate the radiative rates \(R_{ij}\) (both continuum and lines) and the resulting occupation numbers until convergence (details are described in Sect. 3.3).

In total, almost 1000 ALI iterations are required by the complete NLTE procedure, divided into blocks of 30 iterations each. (One iteration comprises calculation of the occupation numbers and the radiation field.) Up to 31 of these iteration blocks are performed using the opacity sampling method (method I), updating the temperature structure and the Rosseland optical depth after each third ALI-iteration, and the total opacities and emissivities after each iteration block. All following iterations are then performed using method II, updating temperature, optical depth, and opacities and emissivities as before, and additionally calculating the blocking factors with the detailed radiative transfer after each iteration block. (Several iteration blocks using method II can be executed, but 1 is usually sufficient – see below.) In this phase the radiative transfer solved in the ALI-iterations within one iteration block is just based on continuum opacities and emissivities, and the blocking factors are applied to get the correct radiative quantities used for calculating the radiative rates.

As a final result of the complete iteration cycle, the converged occupation numbers, the emergent flux, and the final NLTE temperature structure are obtained.

### 3.2. The atomic models

It is obvious that the quality of the calculated occupation numbers and of the synthetic spectrum is directly dependent on the quality of the input data. We have therefore extensively revised and improved the basis of our model calculations, the atomic models.

Up to now, the atomic models of all of the important ions of the 149 ionization stages of the 26 elements considered (H to Zn, apart from Li, Be, B, and Sc) have been replaced in order to improve the quality. This has been done using the Superstructure program (Eissner et al. 1974; Nussbaumer & Storey 1978), which employs the configuration-interaction approximation to determine wave functions and radiative data. The improvements include more energy levels (comprising a total of about 5000 observed levels, where the fine structure levels have been “packed” together2) and transitions (comprising more than 30000 bound-bound transitions for the NLTE calculations and more than 400000 lines for the line-force and blocking calculations3,4), and 20000 individual transition probabilities of low-temperature dielectronic recombination and autoionization.

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2 Note that artificial emission lines may occur in the blocking calculations if the lower levels of a fine structure multiplet are left unpacked but the upper levels of the considered lines are packed – the fine structure levels of an ionization stage should either be all packed or all unpacked.

3 Note further that the consistency of the model calculation requires that the wavelength of the bound-bound transition connecting packed levels in the NLTE calculations to be identical to the wavelength of the strongest component of the multiplet considered in the blocking calculations in order to solve the line radiative transfer and especially the problem of self-shadowing properly.

4 The Superstructure calculations involve many more excited levels than actually used in the NLTE calculation. Our line list does, however, include transitions to such highly excited levels above our limit of considering the level structure; occupation numbers of these upper levels are estimated using the two-level approximation on the basis of the (known) occupation number of the lower level.
### Table 1. Summary of revised atomic data calculated with Superstructure.

In Cols. 2 and 3 the number of levels used in the NLTE calculations are given in packed and unpacked form. Columns 4 and 5 list the number of lines used in the rate equations and for the line-force and blocking calculations.

<table>
<thead>
<tr>
<th>Ion</th>
<th>packed levels</th>
<th>unpacked levels</th>
<th>rate eq. lines</th>
<th>blocking lines</th>
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<td>16 458</td>
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<td>N V</td>
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<td>229</td>
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<tr>
<td>N VI</td>
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<td>S V</td>
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<td>78</td>
<td>404</td>
<td>903</td>
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</table>

Additional line data were taken from the Kurucz (1992) line list: approximately 20 000 lines have been added to the Superstructure data for ions of Mn, Fe, Co, and Ni. These concern transitions to even higher levels than those having been calculated with Superstructure, but which might nonetheless be of significance in the blocking calculations. From the Opacity Project (cf. Seaton et al. 1994; Cunto & Mendoza 1992) another 4 466 lines have been included, as well as photoionization cross-sections (almost 2000 data sets have been incorporated). Collisonal data have become available through the IRON project (see Hummer et al. 1993) – almost 1300 data sets have been included.

Table 1 gives an overview of the ions affected by the improvements. (Users of the program package WM-basic should note that the model calculations will become inconsistent if the atomic data sets are changed haphazardly by those who are not familiar with the source code.)

3.3. The treatment of line blocking

As the thermal width of a UV metal line covers just a few mÅ, a simple straightforward method would require considering approximately $10^7$ frequency points in order to resolve the lines in the spectral range affected by line blocking. Such a procedure would lead to a severe problem concerning the computational time. The alternatives are either to calculate the complete radiative transfer in the comoving frame – again a time-consuming procedure – or to use a tricky method which saves a lot of computation time through the application of some minor approximations (method I), dropping these approximations in the final iteration steps (method II) in order to come to a realistic solution. Our treatment described here uses the second approach.

Although frequently applied, a method using opacity distribution functions (ODFs) (cf. Labs 1951;
Kurucz 1979), where the opacities are rearranged within a rough set of frequency intervals in such a way that a smoothly varying function is obtained which conserves the statistical distribution of the opacities, is not applicable in our case, since there is no appropriate way to treat the lineshift in the wind, and due to the rearrangement of the opacities the frequential position of the lines is changed. This, however, prevents a correct computation of the bound-bound transitions used for the solution of the statistical equilibrium equations.

The approach best suited for our purpose in the first step (method I) is the \textit{opacity sampling} technique (cf. Peytremann 1974; Sneden et al. 1976; Anderson 1991) which compared to the ODF-method is computationally a bit more costly, but does not suffer from the limitations mentioned above. This method allows us to account for the \textit{lineshift} in the wind and the correct influence of line blocking on the bound-bound transitions (cf. Sect. 3, item (ii)), since it preserves the exact frequential position of the lines.

3.3.1. The opacity sampling method (method I)

Following the idea of the \textit{opacity sampling}, a representative set of frequency points is distributed in a logarithmic wavelength scale over the relevant spectral range, and the radiative transfer equation is solved for each point. (For O-stars the actual range depends on \(T_{\text{eff}}\); for hot objects the lower value is at \(\approx 90\) ˚A and for cooler objects the upper value is at 2000 ˚A; note that accurate ionization calculations require extending the line blocking calculations to the range shortward of the \textsc{He}\textsc{II} edge – cf. Pauldrach et al. 1994a.)

In this way the exact solution is reached by increasing the number of frequency points. A smooth transition is obtained when the number of frequency points is increased up to the number \(\approx 10^7\) – which is required to resolve the thermal width of the UV lines. It is obvious however that convergence can be achieved already with significantly less points (see below). Furthermore, special blocking effects on selected bound-bound transitions can be investigated more thoroughly by spreading additional frequency points around the line transition of interest.

In the following subsection we will investigate how many sampling points are required in order to represent the physical situation in a correct way.

The influence of line blocking on the photoionization integrals. The most important effect of line blocking on the emergent spectrum is the influence on the ionization structure via the photoionization integrals

\[
R_{ik} = 4\pi \int \frac{J_\nu}{h\nu} \sigma_{ik}(\nu) \, d\nu. \tag{1}
\]

This can be verified from Fig. 3 where it is shown that the mean radiation field \(J_\nu\) changes rapidly over the frequency interval covered by a typical smooth bound-free cross section \(\sigma_{ik}\) – several 100 ˚A are affected. (Note that dielectronic resonances which may occur in addition are not shown here.)

It is obvious from Fig. 3 that the photoionization rates are sensitive functions of the blocking influence on \(J_\nu\), and hence, on the number of sampling points in the relevant frequency range.

In order to determine the number of sampling points required for an accurate description of the bound-free (photoionization) and bound-bound (line) radiative rates we performed empirical tests by calculating models with an increasing number of sampling points. Representative for the behavior of the radiative rates, and thus the occupation numbers, Fig. 4 shows the dependence of the normalized photoionization integral of the ground state of H in dependence of an increasing number of sampling points within the Lyman continuum.
hydrogen on the number of frequency points. For small numbers of sampling points there is no systematic trend, and the rates converge for higher numbers of sampling points. We conclude that 1000 sampling points within the Lyman continuum on a logarithmic wavelength scale guarantee a sufficient accuracy of about 1 to 2 percent. By means of a separate investigation Sellmaier (1996) showed the given number of sampling points to be reasonable, since it reproduces the actual line-strength distribution function quite well.

The treatment of the lineshift. The total opacity at a certain sampling frequency \( \nu \) is given by adding the line opacity \( \chi_{\text{lb}} \) to the continuum opacity \( \chi_{\text{c}} \)

\[
\chi_{\nu} = \chi_{\text{lb}}(\nu) + \chi_{\text{c}}(\nu),
\]

where \( \chi_{\text{lb}} \) is the sum over all (integrated) single line opacities \( \chi_l \) multiplied by the line profile function \( \varphi_l(\nu) \)

\[
\chi_{\text{lb}}(\nu) = \sum_{\text{lines}} \chi_l \varphi_l(\nu).
\]

Here \( \chi_l \) is

\[
\chi_l = \frac{h\nu_0}{4\pi} (n_i B_{ij} - n_j B_{ji}),
\]

and the analogous expression for the emissivity is

\[
\eta_l = \frac{h\nu_0}{4\pi} n_j A_{ji}.
\]

\((B_{ij}, B_{ji}, \text{and } A_{ji} \text{ are the Einstein coefficients of the line transition at the frequency } \nu_0 \text{, and } h \text{ is Planck’s constant.)}\)

In the static part of the atmosphere a line’s opacity covers its (thermal and microturbulent) Doppler profile \( \varphi_D \) only a very small interval around the transition frequency \( \nu_0 \) (illustrated in Fig. 5 on the right hand side of both figures; note that with regard to our sampling grid about 40 percent of the available lines are treated in this part). The effect of these lines on the radiation field is nevertheless considerable (cf. Fig. 3), if the lines are strong enough to become saturated.

In the expanding atmospheres of hot stars the effect of line blocking is enhanced considerably in the supersonic region due to the nonlinear character of the radiative transfer. A velocity field \( v(r) \) enables the line to block the radiation also at other frequencies \( \nu = \nu_0(1 + v(r)/c) \), i.e., the Doppler shift increases the frequency interval which can be blocked by a single line to a factor of \( \approx 100 \). On the other hand, the velocity field reduces the spatial area where a photon can be absorbed by a line. If a line is optically thick, however, the effect of blocking will ultimately be increased compared to a static photosphere.

The lineshift due to the velocity field is applied to the individual line opacities before the summation in Eq. (3) is carried out at each sampling and depth point (otherwise the effect of the lineshift would be underestimated with respect to the ratio of line width to sampling distance – see below). However, in our approach this is done by applying the Doppler shift of the radial ray to all \( p \)-rays (see Fig. 6), ignoring the angular dependence of the Doppler shift (see below). Apart from the intrinsic character of the sampling method this is the most restrictive approximation in our first iteration cycle. Nevertheless, the main effect of the frequency shift due to the expanding wind – increase of the frequential range of line absorption – is included, and that is what has to be iterated in this first cycle.
From the upper panel of Fig. 5 it is obvious that if the line opacity is simply shifted along the comoving frame frequency (ν_CM) to every radius point successively, many frequency points will miss the line, since the radius grid is too coarse to treat large lineshifts in the observer’s frame. This behaviour is corrected by convolving the intrinsic Doppler profile of the line with a boxcar profile ϕ_D, representing the velocity range around each radius point (Fig. 5, lower panel).

The boxcar profile is the mean profile obtained by considering the velocity shifts Δν of the two corresponding internmesh points (ν_1, ν_2) on both sides of the regarded radius grid point in the way that the gaps in the frequency grid are closed. This can be expressed in terms of the Heaviside function θ:

$$\varphi_{\Delta \nu}(\nu) = \frac{\theta(\nu_2 - \nu) - \theta(\nu_1 - \nu)}{2(\nu_2 - \nu_1)}$$

(6)

ν_1 and ν_2 are the observer’s frame frequencies belonging to the velocities of two successive radius points (r_1 and r_2), i.e., ν_1,2 = ν_0(1 + v(r_1,2)/c). Assuming thermal Doppler broadening for the intrinsic line profile,

$$\varphi_D(\nu) = \frac{e^{-x^2}}{\sqrt{\pi} \Delta \nu_D} \quad \text{with} \quad x = \frac{\nu - \nu_0}{\Delta \nu_D},$$

(7)

where Δν_D is the thermal Doppler width, the convolution (ϕ_D ⊗ ϕ_D)(ν) results in the final profile function

$$\phi(\nu) = (\varphi_D \otimes \varphi_{\Delta \nu})(\nu) = \frac{\text{erf}(x_2 - x) - \text{erf}(x_1 - x)}{2(x_2 - x_1)\Delta \nu_D}. $$

(8)

This profile can be used for the entire sub- and supersonic region. For Δν < ν_therm it gives, as a lower limit, the ordinary opacity sampling, and for sufficiently high velocity gradients (Δv > ν_therm) the integration over a radius interval represents the Sobolev optical depth (τ_{Sob}(r)) of a local resonance zone for a radial ray

$$\Delta \tau = \int_{r_1}^{r_2} \tau_1 \cdot \phi(\nu) \, dr \approx \int_{r_1}^{r_2} \frac{1 - (-1)}{2(x_2 - x_1) \cdot \Delta \nu_D} \cdot (r_2 - r_1)$$

$$= \tau_1 \cdot \frac{r_2 - r_1}{\nu_2 - \nu_1} \cdot \frac{c}{\nu_0} \cdot \left(\frac{d\nu}{dr}\right)^{-1}$$

$$= \tau_{Sob}(r).$$

(9)

At sufficiently high velocity gradients all lines are included in the radiative transfer if the sampling grid is fine enough (see also Sellmaier 1996). In this case our Doppler-spread opacity sampling method therefore becomes an exact solution.

In summary, our Doppler-spread sampling technique makes opacity sampling usable even at large velocity gradients where the standard sampling would miss a line at many frequency points. Broadening of the line with the boxcar profile does not overestimate the line blocking effect, since the convolution (Eq. (8)) preserves the frequency-integrated line strength. Rather, the Sobolev optical depth (Eq. (9)) is the upper limit for the optical thickness of a blocking line as treated with the boxcar profile. The statistical character inherent in opacity sampling is greatly diminished, since at high velocity gradients all available lines are considered.

Furthermore, broadening the lines leads neither to an increased nor a decreased line overlap, since the broadening only spreads a line over the frequencies corresponding to the Doppler shifts between one depth point and the next one. If lines overlap through this broadening at a certain radius grid point, they must also overlap in reality (see Fig. 10) in the interval between that radius point and the adjacent one, because the basic relationship between frequency shift and radius (via the velocity field) is independent of the resolution of the radius grid.

Essentially, the broadening projects the sharply peaked line opacities and emissivities inside a radial interval, which would otherwise be overlooked in the radiation transfer in the discretized scheme, onto a point at the edge of that interval so that the radiation transfer on the discretized radius grid can be performed correctly. For any particular radius interval and spectral line, this affects all frequencies in the interval determined by the Doppler shifts corresponding to the velocities at the edges of the radial interval in question, irrespective of the number of frequency points that actually lie in that frequency interval. Increasing this number of frequency points does not influence this geometric configuration⁵, whereas increasing the radial resolution improves the quality of this procedure, finally converging to the exact solution (cf. method II).

In principle, one would have to account for the angular variation (see Fig. 6) of the Doppler shift in a similar manner, but as no analogy to the boxcar profile method exists in this case, as mentioned above we simply apply the (correctly calculated, radially dependent) opacities (τ_CM(ν, r)) of the central p-ray to the other p-rays, regarding these opacities as being representative. A welcome result of this simplification is the fastness of the method, a very important consideration in this iteration cycle.

Concerning our WM-basic program package running on a normal scalar processor, however, the method is still not fast enough (a model calculation would require an amount of computing time of about 20 hours). The reason is that the Rybicki-method which is used in this step for the solution of the second-order form of the equation of transfer (cf. Mihalas 1978) requires more than 80% of

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⁵ Note that the minimum resolution required for the frequency grid per se is determined by the Doppler spread between the radius grid points, in order to intercept a line at several radius points in succession. As stated above, at sufficiently high velocity gradients all lines are included in the radiative transfer for approximately 1000 frequency points in the Lyman continuum. Moreover, as illustrated in Fig. 4, the chaotic behavior for less than about 1000 frequency points vanishes in this case.
Fig. 7. Logarithm of the ionization fractions of N\textsubscript{iii} (upper panel), N\textsubscript{iv} (middle panel), and N\textsubscript{v} (lower panel) versus density and iteration block number for an O supergiant model ($T_{\text{eff}} = 29\,000\,\text{K}, \log g = 3.0, R_{\ast}/R_{\odot} = 27.0$). The region between two successive contour lines (one iteration block) corresponds to 30 iterations.

In order to illustrate the behaviour of convergence of our method I, the ionization fractions of N\textsubscript{iii}, iv, and v are shown versus density and the iteration block number in Fig. 7 for the first 600 iterations as an example. As displayed, the model converges within 400 iterations – the remaining iterations are required to warrant the luminosity conservation (see Sect. 3.4). The steep increase of N\textsubscript{v} in the wind part results from the EUV and X-ray radiation produced by shock-heated matter (see Sect. 3.5).

We finally note that first results obtained with a version of this procedure as described here so far have already been published. Sellmaier et al. (1996) showed that their NLTE line-blocked O-star wind models solve the long-standing Ne\textsubscript{iii} problem of H\textsubscript{ii}-regions for the first time, and Hummel et al. (1997) carried out NLTE line-blocked models for classical novae.

Special problems. From first test calculations performed in the manner described we recognized and solved two additional nontrivial problems:

The first problem concerns the artificial effect of self-shadowing (see above) which occurs because the incident intensity used for the calculation of a bound-bound transition that enters into the rate equations is already affected by the line transition itself, since the opacity of the line has been used for the computation of the radiative quantities in the previous iteration step. If the lines contained in a frequency interval are of almost similar strength, this is no problem, since the used intensity $I_{\nu}(r)$ calculated at the sampling point represents a fair mean value for the true incident radiation of the individual lines in the interval. If however, a line has a strong opacity with a dominating influence in the interval, the intensity taken at the sampling point for the same bound-bound transition in the radiative rates is much smaller than the true incident radiation for this line, because the line has already influenced this value considerably. In consequence the source function of this line is underestimated and the radiative processes – the scattering part is mostly affected – are not correctly described in the way that the line appears systematically too weak.
The solution to this problem is rather simple: in calculating the bound-bound rates of the dominating lines, we use an incident intensity which is independent of the lines in the considered interval (cf. Pauldrach et al. 1998).

The second problem involves the discretization of the transfer equation in its differential form, for computing the radiative quantities (Feautrier method). In the standard approach (see, for example, Mihalas 1978) the equation of transfer is written as a second-order differential equation with the optical depth \( \tau \) as the independent variable:

\[
\frac{d^2 u}{d\tau^2} = u - S,
\]  

where \( S \) is the source function and \( u = \frac{1}{2}(I^+ + I^-) \), with \( I^+ \) and \( I^- \) being the intensities in positive and negative \( \tau \) direction along the ray considered.

This differential equation is then converted to a set of difference equations, one for each radius point \( i \) on the ray,

\[
\frac{d^2 u}{d\tau^2} \bigg|_{\tau_i} \approx \frac{u_{i+1} - 2u_i + u_{i-1}}{\tau_{i+1}^2 - \tau_{i-1}^2} \tag{11}
\]

resulting in a linear equation system

\[
a_iu_{i-1} + b_iu_i + c_iu_{i+1} = S_i \tag{13}
\]

with coefficients

\[
a_i = -\left(\frac{1}{2}(\tau_{i+1} - \tau_{i-1})(\tau_i - \tau_{i-1})\right)^{-1}, \quad c_i = -\left(\frac{1}{2}(\tau_{i+1} - \tau_{i-1})(\tau_{i+1} - \tau_i)\right)^{-1}, \quad b_i = 1 - a_i - c_i, \tag{14}
\]

(and appropriate boundary conditions). This linear equation system has a tridiagonal structure and can be solved economically by standard linear-algebra means\(^6\). Note that the equations contain only differences in \( \tau \), which can easily be calculated from the opacities and the underlying \( z \)-grid (cf. Fig. 6) as

\[
\tau_{i+1} - \tau_i = \frac{1}{2}(\chi_{i+1} + \chi_i)(\bar{z}_{i+1} - z_i), \tag{15}
\]

with \( \chi_i \) being the opacity at depth point \( i \).

The equation systems are well-behaved if the opacities and source functions vary only slowly with \( z \). Caution must be taken if this cannot be guaranteed, for example, whenever a velocity field is involved at strong ionization edges or with the opacity sampling method at strong lines, since the velocity field shifts the lines in frequency, causing large variations of the opacity from depth point to depth point for a given frequency. In particular, a problematic condition occurs if a point with a larger-than-average source function \( S_i \) and low opacity \( \chi_i \) borders a point with a high opacity \( \chi_{i+1} \) (and low or average source function \( S_{i+1} \)). In reality, this large source function should have little impact, since it occurs in a region of low opacity, and thus the emissivity is small. However, the structure of the equations is such that the emission is computed to be on the order of

\[
\Delta I \approx \mathbf{S} \cdot \Delta \tau \tag{16}
\]

where, if the other quantities are comparatively small (in accordance with our assumptions), the term \( S_i\chi_{i+1} \) dominates\(^7\), leading to artificially enhanced emission.

\(^6\) In practice, a Rybicki-type scheme (cf. Mihalas 1978; and Appendix A, this paper) is used for solving the equation systems for all p-rays simultaneously, since the source function contains a scattering term (see Eq. (A.1)) which redistributes the intensity at each radius shell over all rays intersecting that shell.

\(^7\) The physical reason for the failure of the system is that the source function only has meaning relative to its corresponding opacity. Multiplying the source function from one point with the opacity at another point is complete nonsense.
In Fig. 8 (upper panel) we show the exaggerated emission of the strongest spectral lines in the emergent flux of a stellar model computed using this standard discretization, leading to false results. Even a simple example can serve to illustrate this effect, as demonstrated in Appendix B.

However, with a subtle modification of the equation system coefficients the method can nevertheless be salvaged. The subtle point involves writing the transfer equation as an equation not in \( \tau \), but in \( z \) for derivation of the coefficients, since only this formulation treats correctly the \( z \)-dependence of \( \chi \):

\[
\frac{1}{\chi} \frac{d}{dz} \left( \frac{1}{\chi} \frac{du}{dz} \right) = u - S. \tag{17}
\]

(Note that the grid should still be spaced so as to cover \( \tau \) more-or-less uniformly.) Again approximating the differential equation with a system of differences we obtain

\[
\chi_i \left( \frac{d}{dz} \left( \frac{1}{\chi} \frac{du}{dz} \right) \right)_{i+\frac{1}{2}} \approx \frac{1}{\chi_i} \left( \frac{1}{\chi} \frac{du}{dz} \right)_{i+\frac{1}{2}} - \frac{1}{\chi_i} \left( \frac{1}{\chi} \frac{du}{dz} \right)_{i-\frac{1}{2}}, \tag{18}
\]

\[
\approx \frac{1}{\chi_i} \left( \frac{u_{i+1} - u_i}{z_{i+\frac{1}{2}} - z_i} - \frac{u_i - u_{i-1}}{z_{i+\frac{1}{2}} - z_{i-\frac{1}{2}}} \right), \tag{19}
\]

so that

\[
a_i = - \frac{1}{\chi_i} (z_{i+\frac{1}{2}} - z_{i-\frac{1}{2}}) \cdot \nabla \chi_i (z_i - z_{i+1}), \quad c_i = - \frac{1}{\chi_i} (z_{i+\frac{1}{2}} - z_{i-\frac{1}{2}}) \cdot \nabla \chi_i (z_{i+1} - z_i), \tag{20}
\]

\[b_i = 1 - a_i - c_i.\]

Even though these coefficients seem not too different from those of the standard method, their impact on the computed radiation field is significant, as witnessed by the drastic improvement in the emergent flux shown in the lower panel of Fig. 8. The crucial difference in the coefficients is that the first factor in \( a \) and \( c \) now contains only the local opacity. (We naturally make the corresponding changes in the coefficients of the moments equation as well.)

Test calculations have shown that for the second factor in the coefficients the geometric mean (an arithmetic mean on a logarithmic scale)

\[
a_{i+1, i} = \frac{\chi_{i+1} \chi_i}{\sqrt{\chi_{i+1} \chi_i}} \tag{21}
\]
gives good results, as demonstrated in Fig. 9, where the spectrum of a model computed with the opacity sampling method is compared to that of our detailed radiative line transfer, described in the next section. Considering the relative coarseness of the opacity sampling method, and the fact that the detailed line transfer suffers none of the approximations of the sampling method, the agreement is indeed remarkable. Note again that through our single-ray approximation for the sampling opacities (see above), our method I (opacity sampling) cannot produce P Cygni profiles, since the P Cygni emission is a direct result of the different Doppler shifts of a particular spectral line along different rays.

\[\text{Fig. 9. Comparison of the synthetic EUV spectrum (200–1600 Å) of the S-45 supergiant model (}\; T_{\text{eff}} = 45\,000\,\text{K, log } g = 3.6, R/R_\odot = 18.0)\; \text{calculated with the Doppler-spread opacity sampling method (thick line) and the detailed method (thin line) on the same frequency grid. Due to simplifications in our implementation the sampling method cannot produce P Cygni emission; nevertheless it provides an extremely good basis for the final iterations using the detailed method.}\]

3.3.2. The detailed radiative line transfer (method II)

The detailed radiative line transfer (method II), used for the final iterations, is an exact solution of the transfer equation in the observer’s frame, and is completely equivalent to a comoving frame solution. It removes the two most significant simplifications of our opacity sampling method (method I), i.e., it accounts for:

(1) Correct treatment of the angular variation of the opacities;

(2) Spatially resolved line profiles\(^8\) (implying correct treatment of multi-line effects).

Whereas in method I the former is completely ignored, the lack of spatial resolution was already compensated for to a large extent through the use of our Doppler-spread sampling. (Multi-line interaction is partly included in our method I, but without regard for the sign of the Doppler shift (using just that of the central ray), and without regard for the order of the lines along the ray within a radius interval, as the Doppler-spread sampling effectively “maps” the lines to the nearest radius point.)

With all major approximations removed, the biggest shortcoming that remains in method II is that only Doppler broadening is considered for the lines, as Stark broadening has not yet been implemented. However, this is of no relevance for the UV spectra, as it concerns only a few lines of hydrogen and helium in the optical range.

\(^8\) Note that this will not by itself solve the problem of self-shadowing, since that is an intrinsic property of any method using an “incident radiation” in solving for the bound-bound radiative rates with a continuum already affected by the transition being considered. In the iteration cycle using method II we therefore also have to apply our correction for self-shadowing.
frequency range. It will, however, be important for our future planned analysis of the optical H and He lines. (Stark broadening is not considered in the sampling method either, but here this is of minor significance, as all other approximations are much more serious.)

In contrast to our method I, where the symmetry and our assumption of only radially (not angular) dependent Doppler shifts allowed solving the transfer equation for only one quadrant\(^9\), a correct treatment of both red and blue Doppler-shifted line opacities (see Fig. 6) requires a solution in two quadrants\(^9\) (corresponding to, from the observer’s viewpoint, the front and back hemispheres; the rotational symmetry along the line-of-sight is taken care of through the angular integration weights).

The method employed is an adaptation of the one described by Puls & Pauldrach (1990), using an integral formulation of the transfer equation and an adaptive stepping technique which ensures that the optical depth in each step (“microgrid”) does not exceed \(\Delta \tau = 0.3\), so that the radiation transfer in each micro-interval can be approximated to high accuracy by an analytical formula assuming a linear run of opacity and emissivity between the micro-interval endpoints:

\[
I(\tau_0) = \int_{\tau_0}^{\tau_n} S(\tau) e^{-(\tau-\tau_0)} d\tau + I(\tau_n) e^{-(\tau_n-\tau_0)},
\]

\(9\) The 2nd-order differential representation of the transfer equation accounts for both the left- and right-propagating radiation simultaneously, the unknowns being the symmetric averages of the two.

\(10\) A one-quadrant solution is also possible, but requires both a red- and a blue-shifted opacity for each \((p, z)\)-point and separate treatment of the left- and right-directed radiation, thus being equivalent in computational effort to the two-quadrant solution that solves for radiation going in only one direction.

where the integral is performed as a weighted sum on the microgrid

\[
\int_{\tau_0}^{\tau_n} S(\tau) e^{-(\tau-\tau_0)} d\tau = \sum_{i=0}^{n-1} \left( e^{-(\tau_i-\tau_0)} \int_{\tau_i}^{\tau_{i+1}} S(\tau) e^{-(\tau-\tau_i)} d\tau \right),
\]

each “microintegral” being evaluated as

\[
\int_{\tau_i}^{\tau_{i+1}} S(\tau) e^{-(\tau-\tau_i)} d\tau = w_i^{(a)} S(\tau_i) + w_i^{(b)} S(\tau_{i+1})
\]

with weights

\[
w_i^{(a)} = 1 - \frac{1 - e^{-\Delta \tau_i}}{\Delta \tau_i}, \quad w_i^{(b)} = \frac{1 - e^{-\Delta \tau_i}}{\Delta \tau_i} - e^{-\Delta \tau_i}
\]

where \(\Delta \tau_i = \tau_{i+1} - \tau_i\). To accurately account for the variation of the line opacities and emissivities due to the Doppler shift, all line profile functions \(\varphi_1\) (cf. Eq. (3)) are evaluated correctly for the current microgrid-\((z, p)\)-coordinate on the ray, thus effectively resolving individual line profiles. Only the slowly-varying occupation numbers (or equivalently, the integrated, frequency-independent line opacities \(\nu_{CMF}\) and emissivities \(\nu_{1}\)) and the velocity field are interpolated between the regular radius grid points.

Figure 10 depicts schematically the relationship between the Doppler-shifted frequencies of spectral lines (which are constant in the comoving frame) and the observer’s frame frequency for which the radiative transfer is being calculated. The figure also illustrates the line overlap in accelerating, expanding atmospheres: lines clearly separated in the comoving frame (slices parallel to the \((\nu, \chi)\)-plane) overlap in the observer’s frame (slices parallel to the \((z, \chi)\)-plane at \(\nu_{\text{obs}}\)) due to large Doppler shifts many times the intrinsic (thermal and microturbulent) linewidth. The areas shaded in dark gray correspond to

Fig. 10. Diagram (not to scale) illustrating the basic relationship of the rest-frame frequencies of spectral lines (\(\nu_{\text{CMF}}\)) to observer’s frame frequency (\(\nu_{\text{obs}}\)) for one particular (non-core) \(p\)-ray in the spherically symmetric geometry (cf. Fig. 6). Shown are two spectral lines which get shifted across the observer’s frame frequency by the velocity field in the wind. The dots represent the stepping points of the adaptive microgrid used in solving the transfer equation in the detailed radiative line transfer.
the spatially resolved Sobolev resonance zones of the two lines for this particular observer’s frame frequency and $p$-ray. Note that the dimensions are not to scale, i.e., the intrinsic width of the lines, and consequently the thickness in $z$ of the resonance zones, has been greatly exaggerated in relation to the total velocity shift.

All lines whose maximum Doppler shift $\Delta \nu = \pm v_\infty c / c$ puts them in range of the observer’s frame frequency for which the radiative transfer is being calculated are considered for that frequency point. In Fig. 10, these correspond to those lines whose rest frequencies lie in the gray band in the $(\nu, z)$-plane at $z = 0$.

Broadening of the lines includes thermal and microturbulent contributions. For every atomic species and depth point, the correct thermal Doppler broadening based on atomic weight and local temperature is used. The microturbulence we have used here is about 10 km s$^{-1}$ in the photosphere and grows with the outflow velocity to a maximum value of 0.1 $v_\infty$ in the outer regions of the wind. This is generally in good agreement with the observations.

After the occupation numbers have converged in the iteration cycle using method I, one iteration block with method II is usually sufficient for full convergence of the model, as demonstrated by Fig. 11, where the emergent spectrum of our S-30 model after 1 iteration block of method II is compared to the spectrum resulting from 5 iteration blocks.

A high-resolution spectrum is computed for the purpose of comparison with observations (wavelength range usually from 900 to 1600 Å) after full convergence of the model. This spectrum is generated with the same procedure as used for the detailed line blocking calculations. The high-resolution spectrum is then merged with the (usually) lower-resolution blocking flux for the final flux output (Fig. 12).

### 3.4. Line blanketing

Line absorption and emission also has an important effect on the atmospheric temperature structure. The corresponding influence on the radiation balance is usually referred to as line blanketing. The objective now is to calculate an atmospheric temperature stratification which conserves the radiative flux and which treats the impact of the line opacities and emissivities properly. In principle there are three methods for calculating electron temperatures in model atmospheres. The commonly used one is based on the condition of radiative equilibrium. The second one uses a flux correction procedure, and the third one is based on the thermal balance of heating and cooling rates. As the first method has some disadvantages (see below), we use the second and the third method (cf. Pauldrach et al. 1998; for discussions on calculating temperature structures in other models for expanding atmospheres, see the references listed in Sect. 1; especially Drew (1989, 1990) presented a comprehensive discussion of her method). In deeper layers ($\tau_{Ross} > 0.1$) where true absorptive processes dominate we use the flux correction procedure, and the thermal balance is used in the outer part of the expanding atmosphere ($\tau_{Ross} < 0.1$), where scattering processes start to dominate.

**The flux correction procedure.** The idea of this method is straightforward: the local temperature has to be adjusted in such a way that the radiative flux is conserved. This requires, however, that the temperature is the dominant parameter on which the flux depends, and that the effect of a change in temperature on the flux is known. The first condition is certainly the case for $\tau_{Ross} > 0.1$. With regard to the second condition, a law for the temperature structure is required which is controlled by some global parameters that can be adjusted
The temperature structure versus the Rosseland optical depth and the iteration block number for the same O supergiant model as in Fig. 7 (\(T_{\text{eff}} = 29000\) K, \(\log g = 3.0, R_*/R_\odot = 27.0\)).

Using the method of the \textit{NLTE Hopf functions} we thus have to increase the parameter \(q_0\) first by using the flux deviation at \(\tau_{\text{Ross}} \approx 3\). This parameter is updated in the corresponding iteration cycle until the flux is conserved at this depth on the 0.5% level. Afterwards the same is done with the parameter \(q_\infty\) at an optical depth of \(\tau_{\text{Ross}} \approx 0.1\). In case the flux deviation at \(\tau_{\text{Ross}} \approx 3\) becomes larger than 0.5% the parameter \(q_0\) is iterated again with a higher priority. As a last step in this procedure the parameter \(\gamma\) is adjusted in order to conserve the flux at an optical depth of \(\tau_{\text{Ross}} \approx 1\).

The resulting temperature structure and the corresponding flux deviation for an O supergiant model (\(T_{\text{eff}} = 29000\) K, \(\log g = 3.0, R_*/R_\odot = 27.0\)) are shown in Figs. 13 and 14, respectively. As can be inferred from the figure, the flux is conserved for this model with an accuracy of a few percent. (We note that from test calculations where line blocking was treated, but parameters of the NLTE Hopf function were held fixed, thus effectively ignoring blanketing effects, we found a flux deviation which already starts in the inner part \((\tau_{\text{Ross}} < 50)\) and reaches a value of up to 50% at \(\tau_{\text{Ross}} \approx 0.1\).) This clearly shows the importance of blanketing and backwarming effects and the need to include them. As these test calculations have also shown that absorptive line opacities dominate the total opacity down to an optical depth of \(\tau_{\text{Ross}} > 0.1\), the temperature structure is influenced by backwarming effects in the entire atmosphere – cf. Eq. (26).

The thermal balance. In the outer part of the expanding atmosphere \((\tau_{\text{Ross}} < 0.1)\), where scattering processes start to dominate, the effects of the line influence on the temperature structure are more difficult to treat. Of the two possible treatments, calculating for radiative equilibrium in a proper way in order to conserve the flux. The “Hopf function”, usually applied for the grey case, has an appropriate functional dependence which has been adapted to the spherical NLTE case by Santolaya-Rey et al. (1997) recently in a general way. The main characteristic of the Hopf method is that the Rosseland optical depth is the decisive parameter on which the temperature stratification depends. Thus, in deeper layers the temperature structure can be calculated efficiently by using this new concept of the \textit{NLTE Hopf function}:

\[
T^4(r) = T_{\text{eff}}^4 \frac{3}{4} \left( \frac{\tau_{\text{Ross}}}{\tau_{\text{Ross}}^*} \right) \left( q_N(\tilde{\tau}_{\text{Ross}}) + \tau_{\text{Ross}} \right)
\]

where \(\tilde{\tau}_{\text{Ross}}\) is the radial optical depth in the spherical case,

\[
d\tilde{\tau}_{\text{Ross}} = \chi_{\text{Ross}}(r) \frac{2}{R} \right) \, dr,
\]

and

\[
q_N(\tilde{\tau}_{\text{Ross}}) \equiv q_\infty + (q_0 - q_\infty) \exp(-\gamma \tilde{\tau}_{\text{Ross}})
\]

is the spherical NLTE Hopf function, where the parameters \(q_0, q_\infty,\) and \(\gamma\) are fitted to a predefined run of the \(q_N(\tilde{\tau}_{\text{Ross}})\) stratification (cf. Santolaya-Rey et al. 1997).

Test calculations performed with fixed parameters \((q_0, q_\infty, \) and \(\gamma\)) and without metal lines lead to almost identical results for the temperature structures obtained by our code and the completely independently developed code of Santolaya-Rey et al. (cf. Pauldrach et al. 1998). The reliability of the method has been further proven by the resulting flux conservation which turns out to be on the 1% level.

In the next step, \textit{line blocking} has to be \textit{treated consistently}. Although the line processes involved are complex, they always increase the Rosseland optical depth \((\tau_{\text{Ross}})\). In the deeper layers \((\tau_{\text{Ross}} > 0.1)\) this leads directly to an enhancement of the temperature law (backwarming).
or for thermal balance, we have chosen the latter one, as the convergence of the radiative equilibrium method turned out to be problematic since the $\tau_{\text{Ross}}$ values are small in this part, and, hence, most frequency ranges are optically thin. (This has recently also been proven by Kubat et al. 1999, where the corresponding equations of the method are also presented.) In calculating the heating and cooling rates (Hummer & Seaton 1963), all processes that affect the electron temperature have to be included – bound-free transitions (ionization and recombinations), free-free transitions, and inelastic collisions with ions. For the required iterative procedure we make use of a linearized Newton-Raphson method to extrapolate a temperature that balances the heating and cooling rates.

Figure 13 displays for the model above the resulting temperature structure vs. the number of iterations and shows a pronounced bump and a successive decrease of the temperature in the outer atmospheric part. (For an explanation of this effect, see Mihalas 1978 and references therein.) That the mismatch of the heating and cooling rates immediately goes to 0% in the outer part ($\tau_{\text{Ross}} < 0.07$) where they are applied for correcting the temperature structure has already been presented by Pauldrach et al. (1998).

We note at this place that the direct influence of X-rays (see next section) on the temperature structure via thermalization is negligible, since the energy contained is very small ($L_X/L_{\text{bol}} \approx 10^{-7}$). However, X-rays can indirectly influence the temperature by changing the ionization balance — the corresponding change of opacities in the EUV and UV can then affect the temperature via normal radiative processes. This, however, is properly considered in our procedure.

### 3.5. Revised inclusion of EUV and X-ray radiation

The EUV and X-ray radiation produced by cooling zones which originate from the simulation of shock heated matter arising from the non-stationary, unstable behavior of radiation driven winds (see Lucy & Solomon 1970, who found that radiation driven winds are inherently unstable, and Lucy & White 1980 and Lucy 1982, who explained the X-rays by radiative losses of post-shock regions where the shocks are pushed by the non-stationary features) is, together with K-shell absorption, included in our radiative transfer. The primary effect of the EUV and X-ray radiation is its influence on the ionization equilibrium with regard to high ionization stages like NV and O VI (cf. the problem of “superionization”, the detection of the resonance lines of O VI, N V, S VI in stellar wind spectra (cf. Snow & Morton 1976); in a first step this problem was investigated theoretically by Cassinelli & Olson 1979) where the contribution of enhanced direct photoionization due to the EUV shock radiation is as important as the effects of Auger-ionization caused by the soft X-ray radiation (cf. Pauldrach 1987; Pauldrach et al. 1994a and 1994b). In order to treat this mechanism accurately it is obviously important to describe the radiation from the shock instabilities in the stellar wind flow properly. Note that in most cases a small fraction of this radiation leaves the stellar wind to be observed as soft X-rays with $L_X/L_{\text{bol}} \approx 10^{-7}$ (cf. Chlebowski et al. 1989). Thus, the reliability of the shock description can be further demonstrated by a comparison to X-ray observations, by ROSAT for instance.

In principle, a correct calculation of the creation and development of the shocks is required for the solution of the problem. This means that a detailed theoretical investigation of time-dependent radiation hydrodynamics has to be performed (for exemplary calculations see Owocki et al. 1988; Feldmeier 1995). However, these calculations favor the picture of a stationary “cool wind” with embedded randomly distributed shocks where the shock distance is much larger than the shock cooling length in the accelerating part of the wind. They also indicate that only a small amount of high velocity material appears with a filling factor not much larger than $f \approx 10^{-2}$, and jump velocities of about $u_s = 300 \ldots 700 \text{ km s}^{-1}$ which give immediate post-shock temperatures of approximately $T_s = 1 \times 10^6 \text{ K}$. We also note that the reliability of these results was already demonstrated by a comparison to ROSAT-observations (cf. Feldmeier et al. 1997).

On the basis of these results we had developed an empirical approximative description of the EUV and X-ray radiation, where the shock emission coefficient $\epsilon_\nu(r) = \frac{f}{4\pi} n_p n_e \Lambda_\nu(T_s(r) n_e)$ was incorporated in dependence of the volume emission coefficient $\Lambda_\nu$ calculated by using the Raymond & Smith (1977) code for the X-ray plasma, the velocity-dependent post shock temperatures $T_s$, and the filling factor $f$ which enter as fit parameters — these values are determined from a comparison of the calculated and observed ROSAT “spectrum”. With this description the effects on the high ionization stages (NV, O VI) lead to synthetic spectral lines which reproduce the observations almost perfectly (cf. Pauldrach et al. 1994a and 1994b). However, with this method we were not able to reproduce the ROSAT-observations with the same model parameters simultaneously (see below). We therefore had to determine the filling factor and the post-shock temperatures by a separate and hence in view of our concept not consistent procedure (cf. Hillier et al. 1993). In order to overcome this problem refinements to our method are obviously required.

In the present treatment the outlined approximative description of the EUV and X-ray radiation has been revised. The major improvement consists of the consideration of cooling zones of the randomly distributed shocks embedded in the stationary cool component of the wind. Up to now we had assumed, for reasons of simplicity, that the shock emission is mostly characterized by the intermediate post-shock temperature, i.e., we considered non-stratified, isothermal shocks. This, however, neglects the fact that shocks have a cooling structure with a certain
range of temperatures that contribute to the EUV and X-ray spectrum. Our revision comprises two modifications to the shock structure. The first one concerns the inner region of the wind, where the cooling time can be regarded to be shorter than the flow time. Here the approximation of radiative shocks can be applied for the cooling process (cf. Chevalier & Inamura 1982). The second one concerns the outer region, where the stationary terminal velocity is reached, the radiative acceleration is negligible, and the flow time is therefore large. Here radiative cooling of the shocks is of minor importance and the cooling process can be approximated by adiabatic expansion (cf. Simon & Axford 1966, who investigated a pair of reverse and forward shocks that propagate through an ambient medium under these circumstances). For our purpose we followed directly the modified concept of isothermal wind shocks presented recently by Feldmeier et al. (1997).

Compared to Eq. (29) we account for the density and temperature stratification in the shock cooling layer by replacing the values of the volume emission coefficient \( \Lambda_v(T_s(r) n_e) \) through adequate integrals over the cooling zones denoted by \( \Lambda_v(T_s(r)) \). Thus, \( e_v^s(r) \) is replaced by

\[
e_v^s(r) = \frac{1}{4\pi} n_p n_e \Lambda_v(T_s(r)),
\]

(30)

where

\[
\Lambda_v(T_s(r)) = \pm \frac{1}{x_s} \int_{r}^{r_s} \hat{f}^2(r') \Lambda_v(T_s(r') \cdot \hat{g}(r')) \, dr',
\]

(31)

and \( r \) is the location of the shock front, \( r' \) is the cooling length coordinate with a maximum value of \( x_s \), the plus sign corresponds to forward and the minus sign to reverse shocks, and \( \hat{f}(r') \) and \( \hat{g}(r') \) denote the normalized density and temperature structures with respect to the shock front. The improvement of our treatment is now obviously directly connected to the description of the latter functions. In the present step we used the analytical approximations presented by Feldmeier et al. (1997), which are based on the two limiting cases of radiative and adiabatic cooling layers behind shock fronts (see above).

3.5.1. Test calculations and first results

In the following we present results of test calculations showing the influence of our modified treatment of shock emission. For this purpose we selected the O4f-star \( \zeta \) Puppis as a test object and ignored for the corresponding model calculations the improved blocking and blanketing treatment discussed above. This restriction makes our results directly comparable to those of Pauldrach et al. (1994b), who used the old, simplified treatment for the shock emission. The stellar parameters of \( \zeta \) Puppis, used as basic input for our models, have been adopted from Pauldrach et al. 1994a (see Table 2), together with the abundances listed in Table 3. (These abundances have recently been confirmed by Kahl et al. 2000 and)

Table 2. The stellar parameters of the O4f-star \( \zeta \) Puppis.

<table>
<thead>
<tr>
<th>( \log(\frac{L}{L_{\odot}}) )</th>
<th>( \frac{T_{\text{eff}}}{\text{K}} )</th>
<th>( \log g )</th>
<th>( \frac{\rho}{\rho_{\odot}} )</th>
<th>( v_{\infty}\text{in km/s} )</th>
<th>( 10^{-5} \frac{M}{M_{\odot}/yr} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.006</td>
<td>42</td>
<td>3.625</td>
<td>19</td>
<td>2250</td>
<td>5.9</td>
</tr>
</tbody>
</table>

Table 3. Abundances used for the \( \zeta \) Puppis model. \( Y_X := n_X/n_H \) where \( Y_{X,\odot} \) denotes the solar abundance. For all other abundances solar values were used.

\[
Y_{\text{He}} = 1.20 Y_{\text{He,\odot}} \quad Y_{\text{C}} = 0.35 Y_{\text{C,\odot}} \\
Y_{\text{N}} = 8.00 Y_{\text{N,\odot}} \quad Y_{\text{O}} = 0.75 Y_{\text{O,\odot}}
\]

Table 4. Parameters required for describing the cooling zones of the shocked gas.

<table>
<thead>
<tr>
<th>( \log(\frac{T_s}{\text{K}}) )</th>
<th>( f )</th>
<th>( \log(\frac{N_X}{\text{cm}^{-2}}) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.75</td>
<td>4.3</td>
<td>20.00</td>
</tr>
</tbody>
</table>
from high resolution X-ray spectroscopy of ζ Puppis with the XMM Newton reflection grating spectrometer.)

Although the final objective of our treatment is the determination of the maximum post-shock temperature ($T_s$) and the filling factor ($f$) from a comparison of the calculated and observed ROSAT spectrum, we have also adopted these values for the present test calculations from the similar fits performed by Feldmeier et al. (1997). The values are given in Table 4 together with the interstellar column density of hydrogen (log($N_H$), cf. Shull & van Steenberg 1985).

We start with a spectrum synthesis calculation where EUV and X-ray radiation by shock heated matter is neglected. The comparison between the observed and the synthetic spectrum (Fig. 15) shows clearly that the strong observed resonance lines of O vi are not reproduced by the model. This striking discrepancy illustrates what is meant by the problem of “superionization”. In Fig. 16 we demonstrate, however, that this problem has already been solved by making use of the EUV and X-ray radiation resulting from the treatment of isothermal shocks (model 1). The observed resonance lines of O vi are reproduced quite well, apart from minor differences. Thus it seems that the wind physics are correctly described. That this is not completely the case can be inferred from Fig. 18 where the ROSAT PSPC spectrum (error bars) is shown together with the result of model 1 (thin line). The deficiency of the non-stratified isothermal shocks is obvious – the model yields too little radiation in the soft X-ray part (shortward of 0.7 keV the spectrum is more likely characterized by a cooler shock component of log($T_s$) ≈ 6.3) and too much in the harder energy band.

Following the strategy outlined above we now investigate how far the structured cooling zones behind the shocks can influence this negative result. Figure 18, which shows in addition the calculated X-ray spectrum of our improved model (model 2, thick line), illustrates the improvement. Strikingly, the new calculations can quite well reproduce the ROSAT PSPC spectrum and the comparison shown is at least of the same quality as that obtained by Feldmeier et al. (1997) with their best fit (see also Stock 1998) – note that the total X-ray luminosity of this model is given by

$$L_x = \int_{0.1 \text{ keV}}^{2.5 \text{ keV}} L_\nu \, d\nu = 10^{-7.1} L_{\text{bol}}.$$  \hspace{1cm} (32)

Actually, it is the fact that, compared to the non-stratified isothermal shocks, the post-shock cooling zones with their temperature stratifications radiate much more efficiently in the soft spectral band which leads to the improved fit. This is portrayed in Fig. 19.

Figure 20 shows the location of the optical depth unity in the relevant energy band of ROSAT. Apart from displaying the influence of the K-shell opacities, it becomes evident from this figure that the wind is optically thick up to large radii, especially in the soft X-ray band. This fact reduces the significance of the fit of the ROSAT spectrum, because most of the observed X-ray radiation is obviously
In this section we present the ionizing fluxes and synthetic spectra of a basic model grid of O-stars of solar metallicity, as well as the resonance lines of N\textsuperscript{v} and O\textsuperscript{vi}, since due to their P-Cygni structure these lines provide information about the complete wind region, and the properties of the influencing radiation produced in the whole wind region can therefore be analyzed by means of spectral line diagnostics.

Hence, for the significance of our modified method it is therefore extremely convincing that the synthetic UV-spectra resulting from model 2 also reproduces the observed resonance lines of N\textsuperscript{v} and O\textsuperscript{vi} (the latter is shown in Fig. 17). That both model 1 and model 2 yield a good fit of the P-Cygni lines shows, on the other hand, that distinguishing between two different models from the profiles alone is not always possible. The fact that our improved treatment accounting for the structured cooling zones behind the shocks solves not only the problem of “superionization”, but reproduces for the first time consistently the resonance line spectra of H\textsuperscript{ii}-regions and Planetary Nebulae, and to prove our method and demonstrate its reliability by means of synthetic UV spectra which are qualitatively compared to corresponding observations. (Note that for the standard model calculations the EUV and X-ray shock radiation is not included – using our WM-basic program package this should always be the first step. For succeeding models in an advanced stage we have used solely our previous method based on isothermal shocks (cf. Sect. 3.5), since this is the method which is presently available for WM-basic and thus the models presented in the following can be reproduced by this offered tool.)

Finally, one of the grid models is chosen for a detailed comparison between observed and calculated synthetic spectra, where the primary objective has been to develop diagnostic tools for the verification of stellar parameters, and the determination of abundances and stellar wind properties entirely from the UV spectra. This has been carried out for a cooler O9.5 Ia supergiant, α Cam – a cooler object has been chosen since several aspects tend to make these generally more problematic, such as the ionization balance (more stages are affected) and the optical thickness of the continuum in the wind part.

### 4. Results

#### 4.1. The basic model grid

In this section we present the ionizing fluxes and synthetic spectra of a basic model grid of O-stars of solar metallicity,
comprising dwarfs and supergiants with effective temperatures ranging from 30 000 to 50 000 K. The model parameters, summarized in Table 5, were chosen in accordance with the range of values deduced from observations as tabulated by Puls et al. (1996).

In Fig. 21 we show for each model the primary result, the ionizing emergent flux together with the corresponding continuum. It can be verified from the figure that the influence of the line opacities, i.e., the difference between the continuum and the total flux, increases from dwarfs to supergiants and from cooler to hotter effective temperatures. Both points are not surprising, because they are directly coupled to the mass loss rate (M) which increases exactly in the same manner (cf. Table 5). Due to the increasing M the optical depth of the lines also increases in the wind part and in consequence the line blocking effect is more pronounced. This behavior, however, saturates for objects with effective temperatures larger than T_{eff} = 45 000 K, since in this case higher main ionization stages are encountered (e.g., Fe v and Fe vi) which are known to have

Fig. 21. Calculated ionizing fluxes (Eddington flux in cgs units) versus wavelength of the model grid stars; dwarfs on the left, supergiants on the right.
less bound-bound transitions (cf. Pauldrach 1987). Thus, as can be verified from Fig. 21 the effect of line blocking is strongest for supergiants of intermediate $T_{\text{eff}}$. In Table 5 we also present the numerical values of the integrals of ionizing photons emitted per second for H ($\log Q_{\text{H}}$) and He II ($\log Q_{\text{He}^+}$), as well as the flux at the reference wavelength $\lambda = 5480$ Å, which can be used directly to calculate Zanstra ratios and Strömgren radii.

The next step is to demonstrate the reliability of the calculated emergent fluxes. As the wavelength region shortward of the Lyman edge usually cannot be observed and thus a direct comparison of the fluxes with observations is not possible, an indirect method to test their accuracy is needed. In principle, two such methods exist. The first one is to test the ionizing fluxes by means of their influence on the emission lines of gaseous nebulae, i.e., using the ionizing fluxes as input for nebular models and comparing the calculated emission line strengths to observed ones. However, as a first step this procedure is questionable, since the diagnostics of gaseous nebulae is still not free from uncertainties – dust clumps, complex geometric structure, etc. – and therefore, if
Table 6. Model grid stars and real-world examples with similar spectral types. The parameters of the observed example stars are from Puls et al. (1996).

<table>
<thead>
<tr>
<th>Model / Example</th>
<th>classification</th>
<th>$T_{\text{eff}}$ (K)</th>
<th>log $g$ (cgs)</th>
<th>$R$ ($R_\odot$)</th>
<th>$M$ ($M_\odot$)</th>
<th>log $L$/$L_\odot$</th>
<th>$v_\infty$ (km s$^{-1}$)</th>
<th>$M$ (10$^{-6}$ $M_\odot$/yr)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dwarfs</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D-30</td>
<td></td>
<td>30 000</td>
<td>3.85</td>
<td>8.5</td>
<td>12</td>
<td>37</td>
<td>5.02</td>
<td>1800</td>
</tr>
<tr>
<td>HD 149757 (ζ Oph)</td>
<td>O9 III</td>
<td>32 500</td>
<td>3.85</td>
<td>12.9</td>
<td>43</td>
<td>5.22</td>
<td>1550</td>
<td>≤0.03</td>
</tr>
<tr>
<td>D-40</td>
<td></td>
<td>40 000</td>
<td>3.75</td>
<td>10</td>
<td>21</td>
<td>5.36</td>
<td>2400</td>
<td>0.24</td>
</tr>
<tr>
<td>HD 217068</td>
<td>O7 V n</td>
<td>40 000</td>
<td>3.75</td>
<td>10.3</td>
<td>22</td>
<td>5.39</td>
<td>2550</td>
<td>≤0.2</td>
</tr>
<tr>
<td>D-50</td>
<td></td>
<td>50 000</td>
<td>4.00</td>
<td>12</td>
<td>53</td>
<td>5.91</td>
<td>3200</td>
<td>5.6</td>
</tr>
<tr>
<td>HD 93250</td>
<td>O3 V ((f))</td>
<td>50 500</td>
<td>4.00</td>
<td>18</td>
<td>118</td>
<td>6.28</td>
<td>3250</td>
<td>4.9</td>
</tr>
<tr>
<td><strong>Supergiants</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S-30</td>
<td></td>
<td>30 000</td>
<td>3.00</td>
<td>27</td>
<td>27</td>
<td>5.73</td>
<td>1500</td>
<td>5.0</td>
</tr>
<tr>
<td>HD 30614 (α Cam)</td>
<td>O9.5 Ia</td>
<td>30 000</td>
<td>3.00</td>
<td>29</td>
<td>31</td>
<td>5.79</td>
<td>1550</td>
<td>5.2</td>
</tr>
<tr>
<td>S-40</td>
<td></td>
<td>40 000</td>
<td>3.60</td>
<td>19</td>
<td>53</td>
<td>5.92</td>
<td>2200</td>
<td>10</td>
</tr>
<tr>
<td>HD 66811 (ζ Pup)</td>
<td>O4 I (f)</td>
<td>42 000</td>
<td>3.60</td>
<td>19</td>
<td>53</td>
<td>6.00</td>
<td>2250</td>
<td>5.9</td>
</tr>
<tr>
<td>S-50</td>
<td></td>
<td>50 000</td>
<td>3.90</td>
<td>20</td>
<td>116</td>
<td>6.35</td>
<td>3200</td>
<td>24</td>
</tr>
<tr>
<td>HD 93129A</td>
<td>O3 I f*</td>
<td>50 500</td>
<td>3.95</td>
<td>20</td>
<td>130</td>
<td>6.37</td>
<td>3200</td>
<td>22</td>
</tr>
</tbody>
</table>

Discrepancies are encountered, it is difficult to decide which of the assumptions is responsible for the disagreement. (As an example we mention the Ne iii problem discussed comprehensively by Rubin et al. 1991; Sellmaier et al. 1996.) Rather, nebular modelling and diagnostics should be able to build upon the reliability of the ionizing fluxes, and thus the quantitative accuracy of the fluxes needs to be tested independently of their use in nebular emission line analysis.

The second – and in the light of the difficulties discussed above, the only trustworthy – method is quite analogous, but instead of an external nebula involves the atmosphere of the star itself. The rationale is that the emergent flux is but the outer value of a radiation field calculated selfconsistently throughout the entire wind, which influences the ionization balance at all depths. This ionization balance can be traced reliably through the strength and structure of the wind lines formed everywhere in the atmosphere. Hence it is a natural and important step to test the quality of the ionizing fluxes by virtue of their direct product: the UV spectra of O stars.

4.2. Qualitative comparison with observations

The test is performed by means of synthetic UV spectra which are qualitatively compared to observed IUE spectra and as such cannot be expected to conform in all details. (Minor discrepancies can only be discussed at hand of detailed comparisons. How such discrepancies might be removed by an adjustment of the stellar parameters is shown, as an example, by means of a detailed comparison for α Cam in Sect. 4.3. In several spectral regions with numerous absorption lines (Fe iv, Fe v, Ni iv) discrepancies might also be due to difficulties in placing the continuum of the observations.) For this qualitative comparison we have chosen, for each model of a subset of our grid, a real object from the list of Puls et al. (1996) whose supposed stellar and wind parameters come very close to those of the model. The parameters of the model stars and the selected real objects are summarized in Table 6. (The influence of shock radiation on the models has been neglected at this qualitative step.)

First we investigate the spectra of the dwarf models. As can be inspected from Fig. 22 the comparison of the models D-30 and D-40 with their counterparts HD 149757 and HD 217068 show in principle an overall agreement, whereas the D-50 model, compared with its counterpart HD 93250, shows a severe discrepancy concerning the O v subordinate line at 1371 Å (the calculated line is much too strong) and a less pronounced discrepancy of the N iv subordinate line at 1718 Å (the calculated line is somewhat too weak). Hence we have to realize that either the wind physics is not completely described, or the stellar or the wind parameters of this model are too different from those of HD 93250.

Regarding the first point one might speculate that the inclusion of shock radiation leads to an improvement for the O v line, although this effect would weaken the N iv line further. As is shown below, shock radiation cannot solve the problem, as it does not affect the strength of the O v line at all (cf. the discussion of the S-50 model below). Regarding the second point there are three parameters which could lead to an improvement for both lines. The first one is the effective temperature which, however, would have to be decreased by at least 5000 K. This is on the one hand extremely unrealistic, since O3 and O4 stars would have to be decreased by at least 5000K. This is on the other hand produce another discrepancy due to an increase of the strength of the O iv line at 1338 Å (cf. the S-40 model in Fig. 23). The second parameter is the mass loss rate and the third one is the abundance. In order to investigate whether a systematic variation in the mass loss rate
can solve the problem we computed a small model subgrid for this object by changing the mass loss rate for model D-50, keeping all other parameters the same (cf. Table 7).

The synthetic spectra obtained for the mass loss rate sequence are shown in Fig. 24. As can be seen, lowering the mass loss rate does not solve the problem of the too-strong O\textsc{v} line, as this is one of the last lines to disappear with diminishing mass loss rate, whereas the N\textsc{iv} line which was already too weak disappears immediately. A mass loss rate as low as $10^{-8} M_\odot$/yr would be required to reduce the strength of the O\textsc{v} line to the observed case. The only model where the situation has clearly improved regarding both the N\textsc{iv} and the O\textsc{v} line is D-50-a, the one with a higher mass loss rate. Hence we conclude that the mass loss rate has to be larger by a factor of $\approx 2$ – note that this is also indicated by the stellar parameters of HD 93250 which are very close to those of HD 93129A, which means that the wind parameters have to be similar too; note further the strong similarity of the observed spectra of HD 93129A and HD 93250, which points in the same

Fig. 22. Calculated UV spectra of the model grid dwarfs (left) compared to observed IUE spectra (right) of stars of similar spectral type.
direction (cf. Fig. 25, upper panel); in addition, the abundances of the CNO elements have to be reduced strongly compared to the solar values (for oxygen a reduced value of approximately a factor of 50 is required to weaken the strength of both the O\textsc{iv} and the O\textsc{v} line, as has been inferred from additional test calculations). This, however, should not be the case for the heavier elements (Fe, Ni) since a solar-like abundance of these elements is required to account for the radiative acceleration necessary to produce the higher mass loss rate (cf. Pauldrach 1987).

Now we inspect the comparison of the supergiant models (Fig. 23). As is shown, the observed spectra are reproduced in principle quite well apart from minor differences which can be attributed to a change of abundances (note that the discrepancy of the N\textsc{v} resonance line is due to the omitted shock radiation – see below). Again, the most conspicuous difference regards the O\textsc{iv} and O\textsc{v} subordinate lines which are both too strong, especially for the S-40 and the S-50 models. From the investigation above it is already quite clear that the abundance(s) of

**Fig. 23.** Calculated UV spectra of the model grid supergiants (left) compared to observed IUE spectra (right) of stars of similar spectral type.
the (CN)O element(s) has (have) to be reduced in order to overcome this discrepancy. Nevertheless, we investigate now whether the inclusion of shock radiation leads to an improvement. We have therefore computed an additional model for the S-50 supergiant, model S-50-a, where the influence of shocks on the spectrum has been accounted for; the shock parameters are given in Table 8.

Figure 25 shows that the influence of the shocks on the strength of the O IV and O V lines, and hence the ionization balance, is negligible – just O VI is enhanced selectively, which can be seen by the strength of the O VI resonance line. Another line which is considerably affected by shock emission is the Si IV resonance line. This is because the soft X-ray radiation field of the shocks enhances the ionization of Si V, and thus the recombination to Si IV is decreased (cf. Pauldrach et al. 1994a). As can be inferred from Fig. 25 this improves the fit of the Si IV line significantly. Concerning oxygen we come to the same conclusion as for the dwarf models, namely that the abundance has to be reduced strongly compared to the solar value.

4.3. Detailed analysis of α Cam

In this section we provide, using our S-30 grid model as a starting point, a detailed determination of the abundances and stellar wind properties, and verification of the stellar parameters of α Cam. Special emphasis is given to the shock radiation needed for a fit of the N V and O VI
resonance lines. Although preliminary results from our new shock description look very promising (see Sect. 3.5), this new method is not yet fully implemented in WM-basic. For this investigation we therefore use the method based on isothermal shocks. See Pauldrach et al. (1994a) and (1994b) for a detailed explanation of the shock parameters and the rationale behind the parameterization.

We wish to point out here that we have not attempted an exact determination of \(g\) and stellar radius, but have rather kept the values of our S-3 grid model. The reason is that in contrast to the hydrodynamics the UV spectrum depends only marginally on these parameters, the main influence being due to \(T_{\text{eff}}\) and \(M\) (i.e., density). The radius and the surface gravity \(\log(g)\) can in principle be determined from a selfconsistent calculation of the hydrodynamics and the NLTE model, in which both values would be adapted in such a way that the hydrodynamics, with consistent force multiplier parameters from the NLTE occupation numbers (in turn again dependent on the hydrodynamic solution), would yield the mass loss rate and the terminal velocity deduced from \(H_{\alpha}\) and the observed UV spectrum (cf. Pauldrach et al. 1994a, 1994b). This latter procedure has however not yet been implemented in WM-basic.

In Fig. 26 we compare the spectrum of our basic S-30 grid model with the spectra of \(\alpha\) Cam observed with IUE and Copernicus. With the exception of a few strong lines, notably the NV resonance doublet at 1238, 1242 Å and the subordinate C III line at 1247 Å, the agreement is very good for a first step, especially with respect to the iron and nickel “forest” between 1400 and 1600 Å. As the two lines mentioned above are mainly affected by shock radiation, and as it is not clear a priori how shocks affect the lines used to determine, e.g., the temperature, we will first attempt to fit the shock parameters before verifying the stellar parameters. We note in passing that the strong lines from 1000 to 1100 Å (with the recurring pattern), that complicate an exact fit of the O VI resonance line, are absorption by interstellar molecular hydrogen.

A model with reasonable “first guess” values for the shock parameters (model a, Table 9) already gives a very good fit to the NV resonance line, as shown in Fig. 27, the emission, however, completely decimated by the much too strong C III λ1247 line. Increasing the X-ray luminosity \(\log(L_X/L_{\text{bol}})\) to reduce the C III occupation unfortunately also tends to ionize Si III to Si IV, thus reducing the strength of the Si III line. Modifying \(v_t/v_\infty\) (the ratio of the maximum jump velocity to the terminal velocity, where \(v_t\) characterizes the immediate post-shock temperature; see Pauldrach et al. 1994a) does not change this fact, as we have confirmed by calculating a grid of models with values of \(\log(L_X/L_{\text{bol}})\) ranging from −8 to −6.5 and \(v_t/v_\infty\) ranging from 0.1 to 0.2. For example, a model sequence (models b1–b3) is shown in Fig. 28, in which we vary \(\log(L_X/L_{\text{bol}})\) from −7.5 to −6.5, keeping all other parameters constant (cf. Table 9). As can be seen, the Si III line already begins to weaken, while C III still remains too strong. Adjusting the parameter \(\gamma\) (which controls the strength of the shocks relative to the local velocity – cf. Pauldrach et al. 1994a) does not help either, as both the C III and the Si III line are formed in the inner part of the wind, thus both being subject to the same radiation field.

From this we conclude that solar abundances cannot reproduce the observed spectrum. A model, however, with a reduced carbon abundance of one tenth solar (model c) can indeed give a good fit of the line, as shown in Fig. 29. We will use this abundance for the following calculations, unless stated otherwise. As carbon thus shows indications of the CNO-process, we have for reasons of consistency also increased the nitrogen abundance by a factor of 10, but such a large factor is not compatible with the fit of the nitrogen lines; from the final models we determined the nitrogen abundance to be approximately solar.

Despite the good fit, this model is still not satisfactory, as it shows no signs of O VI. Since the C III/Si III-balance strongly constrains the shock strength in the inner regions, this cannot be achieved simply by increasing the X-ray
luminosity. However, the onset of shocks can be adapted with the parameter $m$, which gives the ratio of outflow to sound velocity where shocks start to form, and as the jump velocity is correlated to the outflow velocity, the corresponding radius (cf. Pauldrach et al. 1994a). Increasing this parameter allows shocks in the outer regions ($O\text{VI}$ appears close to $v_{\infty}$), while leaving the inner regions (where $Si\text{III}$ is present) largely undisturbed. This behaviour of the shocks is already an important result from our analysis.

Figure 30 shows a model sequence in which the radius at which shocks start to form has been successively moved outwards, keeping all other parameters equal. In model d4, where $Si\text{III}$ remains fairly unaffected by shocks, the P Cygni emission of $O\text{VI}$ and $N\text{V}$ has completely disappeared.

Fig. 30. Models d1–d4, a sequence in which the radius at which shocks start to form has been successively moved outwards, keeping all other parameters equal. In model d4, where $Si\text{III}$ remains fairly unaffected by shocks, the P Cygni emission of $O\text{VI}$ and $N\text{V}$ has completely disappeared.

Fig. 31. Model e. Increasing the shock jump velocity provides for harder shock radiation that has a stronger influence on $O\text{VI}$ and $N\text{V}$ than on $Si\text{III}$. The abundances of $C$, $N$, $O$, and $P$ have also been adapted.

This reasoning is confirmed by model e (Fig. 31), in which we have reduced $m$ to 30 and increased $v_{s}/v_{\infty}$ to 0.25 (corresponding to a maximum shock temperature of $2.0 \times 10^{6}$ K). The shock parameters of this model yield the best overall agreement of the most important UV spectral
Table 9. Model parameters for α Cam. Abundances not explicitly mentioned are solar; if given, the number is the factor relative to the solar value. \( T_{\text{eff}} \) is in K, \( \dot{M} \) in \( 10^{-6} M_\odot/\text{yr} \).

<table>
<thead>
<tr>
<th>Model</th>
<th>( T_{\text{eff}} )</th>
<th>( \dot{M} )</th>
<th>( \nu / \nu_\infty )</th>
<th>\log(( L_X / L_{\text{bol}} ))</th>
<th>( \gamma )</th>
<th>( m )</th>
<th>abundances</th>
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<tr>
<td>a</td>
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<td>1</td>
<td>solar</td>
</tr>
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<td>5</td>
<td>0.14</td>
<td>-7.5</td>
<td>1</td>
<td>1</td>
<td>solar</td>
</tr>
<tr>
<td>b2</td>
<td>30 000</td>
<td>5</td>
<td>0.14</td>
<td>-6.0</td>
<td>1</td>
<td>1</td>
<td>solar</td>
</tr>
<tr>
<td>b3</td>
<td>30 000</td>
<td>5</td>
<td>0.14</td>
<td>-6.5</td>
<td>1</td>
<td>1</td>
<td>solar</td>
</tr>
<tr>
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<td>-7.5</td>
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<td>1</td>
<td>C = 0.1, N = 10., O = 1.0, P = 0.1, S = 2.0</td>
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<tr>
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<td>5</td>
<td>0.14</td>
<td>-7.0</td>
<td>0.5</td>
<td>1</td>
<td>---</td>
</tr>
<tr>
<td>d2</td>
<td>30 000</td>
<td>5</td>
<td>0.14</td>
<td>-7.0</td>
<td>0.5</td>
<td>30</td>
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<tr>
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<td>-7.0</td>
<td>0.5</td>
<td>40</td>
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<td>-7.0</td>
<td>0.5</td>
<td>60</td>
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</tr>
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<td>0.25</td>
<td>-6.5</td>
<td>0.5</td>
<td>30</td>
<td>C = 0.1, N = 2.0, O = 0.3, P = 0.05, S = 1.0</td>
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<td>f</td>
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<td>0.25</td>
<td>-6.5</td>
<td>0.5</td>
<td>30</td>
<td>---</td>
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<tr>
<td>h</td>
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<td>5</td>
<td>0.25</td>
<td>-6.5</td>
<td>0.5</td>
<td>30</td>
<td>---</td>
</tr>
<tr>
<td>i</td>
<td>28 000</td>
<td>2.5</td>
<td>0.25</td>
<td>-6.5</td>
<td>0.5</td>
<td>30</td>
<td>---</td>
</tr>
<tr>
<td>j</td>
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<td>0.25</td>
<td>-6.5</td>
<td>0.5</td>
<td>30</td>
<td>---</td>
</tr>
</tbody>
</table>

best fit →

| g     | 29 000          | 5              | 0.25            | -6.5            | 0.5     | 30      | C = 0.05, N = 1.0, O = 0.3, P = 0.05, S = 1.0 |
| h     | 28 000          | 2.5            | 0.25            | -6.5            | 0.5     | 30      | ---           |
| i     | 28 000          | 2.5            | 0.25            | -6.5            | 0.5     | 30      | ---           |
| j     | 28 000          | 10             | 0.25            | -6.5            | 0.5     | 30      | ---           |

Fig. 32. The complete EUV spectrum of model e \( (T_{\text{eff}} = 30\,000 \text{ K}) \).

Fig. 33. Model f. Like model e, but with a \( T_{\text{eff}} \) of 29 000 K. Note the better fit of SiIII and OIV.

lines. Note that for this model we have in addition reduced the phosphorus abundance to 0.05 times solar, to obtain a fit of the P v line. As it concerns only a single element and since evidence for an underabundance of phosphorus has already been encountered from an analysis of another hot star, namely the O4f-star ζ Puppis (cf. Pauldrach et al. 1994a; and Sect. 3.5, this paper), it is most likely that the discrepancy between the observed and calculated P v resonance line is caused by the proposed abundance effect. We further note that the imbalance of Si IV and S IV is most likely due to a combination of abundances and imperfect atomic data, as our Si IV atomic model has not yet reached the quality of those of other ionization stages and is still in a stage of rather incomplete description (cf. Table 1). Note that we have also reduced the oxygen abundance, thereby improving the fit of both O IV and O VI.

Having thus constrained the strength and distribution of the shocks, we still need to check to what extent the effective temperature and the mass loss rate can be constrained further through our analysis of the UV spectrum. For this purpose, we have computed a model with \( T_{\text{eff}} \) of 29 000 K but otherwise same parameters (model f), whose spectrum is shown in Fig. 33. Comparing this to the spectrum of model e \( (30\,000 \text{ K}) \) (Fig. 32) we note a marginally better fit of the spectral region from 1450 to 1650 Å, and a slight improvement in Si IV and O IV. More significant is the extreme sensitivity of the Si III and the C III line to this small change in temperature, which reveals that these lines can be utilized as temperature indicators in this spectral range. We concede that the fit of N V is still somewhat imperfect, but point out that the shock model used is not in a final stage (see above); we expect this to improve with our new method. In a next step we have reduced the carbon abundance further (to 0.05 solar) and nitrogen to solar abundance, as this improved not only the N V/C III fit, but also the N II line at 1183, 1185 Å (model g, Fig. 34). Note also that the saturated C IV
Fig. 34. Model g. Like model f, but with adapted abundances (see Table 9). This is our best fit.

resonance line is not affected by this change of the carbon abundance.

If one assumes the conservation of CNO abundances and believes that the abundances produced in the innermost part of the star appear in the same ratio at the surface of the star, then the results obtained above for the CNO abundances indicate that the ZAMS abundances of the light elements in α Cam deviate from the solar values. (Latest evolutionary calculations by Meynet & Maeder 2000, incorporating the effects of rotational mixing show that processed material is more efficiently transported to the surface than previously thought.) Our analysis shows a nitrogen overabundance relative to carbon and oxygen; whether this will have implications concerning the metal dependence of the formation of massive stars is not clear at this time.

Lowering the effective temperature further to 28 000 K worsens the fit, as the iron and nickel lines around 1500 Å become too strong (Fig. 35). As the strength of these lines depends also on the mass loss rate, it is conceivable that an adaptation of this parameter can again improve the fit. However, $\dot{M}$ is strongly constrained by the strength of the He II line, as demonstrated in Figs. 36 ($\dot{M} = 2.5$) and 37 ($\dot{M} = 10$). We have confirmed in test calculations that the iron abundance must be solar ($Z = 1.0$) to reproduce the observed spectrum and to account for the radiative acceleration needed to produce the observed mass loss rate (cf. Pauldrach 1987), a finding compatible with our earlier statement concerning the relative abundances of iron to the light CNO elements.

Finally, we wish to illustrate the influence of the shocks on the ionizing flux of α Cam. In Fig. 38 we compare

Fig. 35. Model h. Same as model g, but with $T_{\text{eff}} = 28 000$ K.
the ionizing flux of our S-30 supergiant grid model, where shocks have been neglected, to that of our model e for α Cam. (We have used model e for this comparison, and not our final model g, since model e and the grid star S-30 both have the same effective temperature of 30000 K. The other models (a–j) of course have similar X-ray and EUV fluxes to that of model e.) Due to the lower optical depth redward of the He II edge, the shock radiation enhances the flux as far as redward as 400 Å; this might have important implications concerning the solution of the Ne III problem (cf. Sellmaier et al. 1996), pending further investigation. Note: the Zanstra integral for He II is increased from $log Q_{\text{He II}} = 6.39$ for the S-30 model to 15.07 for model e!

(The influence of the shocks on the EUV spectrum is larger in α Cam than in ζ Puppis, due to the lower effective temperature and thus overall lower EUV fluxes – this is in agreement with MacFarlane et al. 1994, who show how the importance of X-rays decreased from B type stars and late O types to earlier types. The influence of the shocks on the whole spectral regime from EUV to radio has been shown by Pauldrach et al. 1994b, their Fig. 11, and especially for EUV and UV by Sellmaier et al. 1993; the expected influence on the ionizing fluxes has also been discussed by Schaerer & de Koter 1997.)

5. Conclusions

After a long period of work in the areas of non-LTE radiative transfer, hydrodynamics, and atomic physics we have now developed a fast numerical model code for expanding atmospheres which incorporates for the first time the required physics without restrictive approximations – rate equations for individual levels of all ions using detailed up-to-date atomic models, the equations of stationary radiation hydrodynamics, the energy equation, the radiative transfer equation including the effects of overlap of numerous spectral lines of different ions, and a realistic description of shock emission from instabilities in the stellar wind flow – thus making a quantitative analysis of observed high-resolution UV spectra via comparison with synthetic spectra reasonable. One of the most complicating effects in this complex physical system is the overlap of thousands of spectral lines of different ions. Especially concerning this latter point we have made significant progress with regard to our model code; the decisive factor has been to relax some rather severe approximations concerning the correct treatment of Doppler-shifted line radiation transport and the corresponding coupling with the radiative rates in the rate equations. We have demonstrated that these modifications to the models concerning the energy distributions, ionizing continua, and line spectra lead to much better agreement with the observed UV spectra. This has important repercussions for the quantitative analysis of hot star spectra.

With this new method in hand we have already presented a new basic model grid of stars of solar metallicity that can be used as input for the analysis of spectra of

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**Fig. 36.** Model i. As model h, but with a lower mass loss rate ($\dot{M} = 2.5 \times 10^{-6} M_\odot$/yr compared to $\dot{M} = 5 \times 10^{-6} M_\odot$/yr for model h).

**Fig. 37.** Model j. As before, but with a higher mass loss rate ($\dot{M} = 10 \times 10^{-6} M_\odot$/yr). Note the change in the strength of the He II line.

**Fig. 38.** Comparison of the ionizing flux of our S-30 grid star, where shocks have been neglected, to that of model e, showing the non-negligible influence of shocks on the ionizing fluxes.
emission line nebulae. The qualitative diagnostic investigation performed on basis of the model grid revealed that for the most massive stars of our sample (HD 93129 A and HD 93250) the oxygen abundance is considerably reduced compared to the solar value. In fact, it is conceivable that these stars are extremely massive precisely because the cooling behavior of the protostellar clouds from which they formed is correlated with a lower oxygen abundance. This would mean, however, that a lower oxygen abundance should be observed in all massive stars younger than O5. This conclusion is in addition supported by the fact that no very strong O v lines are observed for this kind of objects.

From the first detailed analysis of the O9.5 supergiant α Cam we conclude that our spectrum synthesis technique does, in principle, allow the determination of effective temperature and abundances – in fact, a determination of the effective temperature to within ±1000 K and of the abundances to within a factor of 2 seems not unreasonable. Carbon and phosphorus show clear signs of an underabundance on the order of one tenth its solar value, as does oxygen with about 0.3 solar, whereas the abundance of iron must be roughly solar to reproduce the spectrum of the numerous Fe IV and Fe V lines. To produce the ionization balance observed in the lighter elements C, N, O, and Si, the influence of shock radiation must start at larger radii where for shorter wavelengths the largest shock temperatures dominate. Thus, the way the X-ray spectral region selectively affects the ionization balance of different elements, observable through the lines in the EUV spectrum, provides constraints on the lower shock temperatures; we have determined maximum shock temperatures on the order of 2.0 × 10⁶ K. Especially the Si III and C III lines have been found to be invaluable diagnostic instruments for this purpose. Our detailed analysis of the UV spectrum and the shocks needed to reproduce the observed lines has led to a significant difference in the ionizing flux compared to models without shocks. Thus we conclude that this type of analysis is indispensable and must be regarded as the ultimate test for the accuracy of ionizing fluxes from models.

Our research plan for the future has three major objectives. First, we will have to implement further improvements to the model atmosphere code, especially concerning the planned analysis of optical lines. For this purpose Stark broadening has to be included for the purposes (cf. Sellmaier et al. 1993, who found that comoving frame treatment the Sobolev-plus-continuum approximation leads to a non-negligible change of the strength of some H and He lines). In connection with this some minor approximations will also have to be checked in detail.

Second, we plan to apply the model atmosphere code to a comprehensive sample of stars of different metallicities for modelling ionizing fluxes and line spectra, the qualitative review of the model grid presented in this paper being just a first step for a detailed quantitative analysis of ionizing fluxes and quantitative verification of the accuracy and reliability of the models. Furthermore, spectral analysis in the UV and the optical range of individual stellar objects in Local Group galaxies and in galaxies as distant as the Virgo Cluster will be performed.

Third, we plan on using the technique of population synthesis to calculate integrated spectra for a large range of stellar metallicities from synthetic UV spectra of our new models, for the determination of stellar abundances and the physical properties of the most UV-luminous stars in star-forming galaxies even at high redshifts. With our new diagnostic tool of detailed models for expanding atmospheres, and the observations of the HST space observatory and the ESO VLT ground-based telescope which are already available, the concept of using luminous hot stars for quantitative UV spectral analyses for determining the properties of young populations in galaxies is not only reasonable, but first tentative steps in this direction have already been taken (see, for instance, Mehler et al. 2000, 2001, whose first diagnostic investigations have shown that, considering proper reddening, the spectra of galaxies they had observed at high redshifts (z ~ 3) can in principle be fitted with synthetic spectra from our hot star models).

Acknowledgements. We wish to thank our colleagues Dr. J. Puls, Dr. R.-P. Kudritzki and Dr. K. Butler for helpful discussions and Dr. S. Becker for help in improving the atomic data. It is a special pleasure to thank Dr. J. Puls additionally for providing us with a basic solver routine for the formal integral. We also thank the anonymous referee for convincing us of the necessity of making the complexities of our method understandable for everyone. This research was supported by the Deutsche Forschungsgemeinschaft in the “Gerhard Hess Programm” under grant Pa 477/2-3, by the DLR under grant 50 QV 9704 1, and by the German-Israeli Foundation under grant I-551-186.07/97.

Appendix A: A new concept for a fast solution of the Rybicki-method

Here we present for the solution of the Rybicki-scheme a concept which is compared to the standard procedure 10 times faster on a vector processor and 3 to 5 times faster on a scalar processor (see Sect. 3.3).

We start from the final system for the solution of the mean intensity – the vector J describes its depth variation (the number of depth points is N):

\[ J = W^{-1} Q \]  \hspace{1cm} (A.1)

with \( N' \) is the number of p-rays

\[ W = \tilde{W} - 1 = -1 - \sum_{j=1}^{N'} T_j^{-1} U_j, \quad Q = - \sum_{j=1}^{N'} T_j^{-1} K_j \]  \hspace{1cm} (A.2)
where
\[ T_j = T'_j w_j^{-1}, \quad w_j = \begin{pmatrix} w_{1,j} \\ \vdots \\ w_{N_j,j} \end{pmatrix}, \quad (A.3) \]
and \(N_j\) is the number of radius points for the \(j\)th \(p\)-ray (cf. Fig. 6). \(w_j\) is the diagonal matrix of the integration weights, and \(T'_j\) is a tri-diagonal matrix defined by the coefficients of the difference equation of transfer
\[ a_{i,j} u_{i-1,j} + b_{i,j} u_{i,j} + c_{i,j} u_{i+1,j} - \beta_i J_i = (1 - \beta_i) S_i \quad (A.4) \]
where \(a_{i,j}, b_{i,j}, \) and \(c_{i,j}\) are the coefficients as in Eq. (20) (Sect. 3.3.1) for the \(j\)th \(p\)-ray,
\[ T'_j = \begin{pmatrix} b_{1,j} & c_{1,j} \\ \vdots & \ddots & \vdots \\ a_{N_j,j} & b_{N_j,j} \end{pmatrix}. \quad (A.5) \]
The variables \(u_{i,j}\) are the symmetric averages of the intensities, the coefficients \(\beta_i\) are the ratios of Thomson-opacities to total opacities, and the diagonal matrix \(U_j\) and the vector \(K_j\) are defined as
\[ U_j = \begin{pmatrix} -\beta_1 \\ \vdots \\ -\beta_{N_j} \end{pmatrix}, \quad K_j = \begin{pmatrix} (1 - \beta_1) S_1 \\ \vdots \\ (1 - \beta_{N_j}) S_{N_j} \end{pmatrix}. \quad (A.6) \]
The usual, but time-consuming solution method is to calculate the inverse matrices \(T^{-1}_j\) by a forward-elimination and back-substitution procedure and from these, obtaining \(Q\) and \(W\), and, finally, \(J\).

Due to the diagonal character of the matrix \(U_j\) this is, however, not necessary, because from the solution of the first column of \(W\) obtained by using just the first column of \(U_j\) in Eq. (A.2) the solutions of the remaining columns of \(W\) can be generated. Hence, the elimination procedure only has to be applied to the following two sets of equations (\(N'_i\) is the maximum number of \(p\)-rays for the \(i\)th radius point):
\[ Q_i = -\sum_{j=1}^{N'_i} K_{j,i} l \quad (A.7) \]
\[ \tilde{W}_{i,1} = -\sum_{j=1}^{N'_i} \tilde{U}_{j,i} = -\sum_{j=1}^{N'_i} \tilde{B}_{j,i} V_{j,1} \quad (A.8) \]

with
\[ \tilde{K}_j = T^{-1}_j K_j, \quad \tilde{B}_j = T^{-1}_j B_j \quad (A.9) \]
where
\[ V_{j,1} = \begin{pmatrix} -\beta_1 \\ \vdots \\ -\beta_1 \end{pmatrix}, \quad B_j = \begin{pmatrix} 1 \\ 0 \\ \vdots \\ 0 \end{pmatrix}. \quad (A.10) \]

Note that since the solution procedure is not recursive with respect to the index \(j\) the elimination can be performed simultaneously for all \(p\)-rays. Note further that the structure of the sums in Eqs. (A.7) and (A.8) are equivalent to the operations \(x = A \cdot y\) where \(A\) is a triangular matrix. Thus, BLAS level-2 routines can be applied.

For the construction of the remaining columns of \(W\) we now make use of the already calculated matrix \(\tilde{B}_{j,1}\) and two auxiliary matrices obtained as a byproduct during the forward-elimination procedure by solving Eq. (A.8).

The first one is:
\[ F'_{j,i} = F'_{j,i-1} f_{j,i}, \quad i = 1, \ldots, N - 1 \quad (A.11) \]
with
\[ F'_{j,0} = 1, \quad f_{j,i} = -a_{j,i+1} / b_{j,i}. \quad (A.12) \]
Here \(b_{j,i}\) is the updated value of \(b_{j,i}\) obtained after the forward-elimination step by solving Eq. (A.8) \(b_{j,i+1} = b_{j,i+1} + f_{j,i} c_{j,i}\).

The second one is:
\[ F''_{j,i} = F''_{j,i+1} + g_{j,i}, \quad i = N - 1, \ldots, 1 \quad (A.13) \]
with
\[ F''_{j,N} = 1, \quad F''_{N-1,N} = 1, \quad g_{j,i} = -c_{j,i} / b_{j,i}. \quad (A.14) \]
The final step – construction of the columns \(l = 2, \ldots, N\) of \(W\) (where \(l\) denotes the position of the unity value for the vector \(B_j\) in Eq. (A.10) – consists of two parts, where the first part replaces the forward-elimination and the second part the back-substitution procedure:

In the first part the components \(i = l, \ldots, N\) of the \(l\)th column of \(\tilde{W}\) are determined by
\[ \tilde{W}_{i,l} = -\sum_{j=1}^{N'_i} \tilde{B}_{j,i} V_{j,l} \quad (A.15) \]

where
\[ V_{j,l} = V_{j,l} / F'_{j,l-1}. \quad (A.16) \]
Again, a triangular matrix has simply to be multiplied by a vector using a BLAS level-2 routine.

In the second part the components \(i = l - 1, \ldots, 1\) of the \(l\)th column of \(W\) are determined by
\[ \tilde{W}_{i,l} = -\sum_{j=1}^{N'_i} F''_{j,l} V''_{j,l} \quad (A.17) \]
where
\[ V''_{j,l} = V''_{j,l} / F''_{j,l}. \quad (A.18) \]
Now, a rectangular matrix has simply to be multiplied by a vector again using a BLAS level-2 routine. Hence, instead of a number of operations \(\sim N^3\) a number of operations only \(\sim N^2\) must be performed, and as extremely fast routines are used to solve the non-recursive system, the solution of the Rybicki-scheme is now almost as fast as the solution of the moments equation.
Appendix B: A simple demonstration of the failure of the standard \( \tau \) discretization of the equation of transfer

Let us assume a one-dimensional, sharply bounded cloud of material of moderate opacity,

\[
\chi = \begin{cases} 
1, & \text{if } |z| < 5 \\
10^{-10}, & \text{if } |z| \geq 5 
\end{cases}
\]

with a symmetric boundary condition at \( z = 0 \) (the middle of the cloud) and no influx from the outside (i.e., \( I^- = 0 \) at \( z = 10 \)).

With a constant source function \( S = 1 \) everywhere, both the old and the new discretization of the transfer equation yield essentially the same radiation field (upper panel of Fig. B.1), a solution which is immediately obvious: the center of the cloud (\( z = 0 \)) is optically thick, the intensity there thus equal to the source function. Towards the edge of the cloud, more and more radiation escapes, and the intensity decreases. On the outside of the cloud, essentially no emission is produced (nothing is absorbed, either); only the radiation emitted by the cloud contributes to the local intensity (since we have no influx from the right), therefore \( u = \frac{1}{2} S \).

The results change considerably if we adopt a high source function of \( S = 100 \) for the single grid point at \( z = 5 \), the first grid point outside of the cloud. As illustrated in the lower panel of Fig. B.1, the old method produces extra emission on a scale of

\[
\Delta I = \frac{1}{2} S \cdot \Delta \tau \approx \frac{1}{2} S \cdot 100 \cdot 10^{-10} \Delta z
\]

\[
\approx \frac{1}{4} 100 \Delta z.
\]

For the grid with 100 points, \( \Delta z = 0.1 \), giving us an intensity

\[
u \approx 0.5 + \frac{1}{2} \cdot \frac{1}{4} 100 \cdot 0.1 = 1.75
\]

outside the cloud. (0.5 is from the normal emission of the cloud as before, and the factor 1/2 in the second term is again due to the fact that \( u = \frac{1}{2} (I^+ + I^-) \), with \( I^- = 0 \) on the outside of the cloud, so that \( u = \frac{1}{2} I^+ \).) The extra emission is proportionally reduced in the finer grids as \( \Delta z \) gets smaller.

The new method, on the other hand, is essentially indifferent to the grid spacing in this particular configuration, since the emission is only computed on the basis of the local opacity, which is low for the point in question. If we were to choose a point inside the cloud to have a high source function, then both methods again produce similar results, also dependent on grid spacing, since in this case the point in question does have a high opacity, and therefore produces substantial emission proportional to the interval length.

Concluding, it can be said that the discretization methods differ in the assumptions that are made about the emissivity (in the old method, based on average values of opacity and source function assuming linear run between the grid points; in the new method, extrapolating the local value) – information not available given just the values at the grid points, and which must therefore be supplied separately through the choice of discretization coefficients. It just so happens that the new method produces results which are more compatible with our detailed formal integral.

References
