# Documentation

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## **Contents**

1	Introduction	3
	1.1 Radiation hydrodynamics	3
	1.2 Radiative transfer	3
	1.3 Philosophy	3
2	Installation	4
_	2.1 Requirements	4
	2.2 Getting the code	4
	2.3 Quick start	4
	2.4 Installing HDF5	4
	2.5 Importing the plotting routines	5
3	Main code modules	5
	3.1 line3D	5
	3.1.1 Program model.eo	6
	3.1.2 Program sc3d.eo	8
	3.1.3 Program modelspec.eo	9
	3.1.4 Program spec.eo	10
	3.2 boss3D	11
	3.2.1 The namelist files	12
	3.3 cont3Dslab	14
4	Getting started	14
5	Related papers	14
6	Cite	14
7	Developers and contributors	14
8	Known problems and solutions	14
o	•	
9	ТоДо	14
10	Acknowledgements	15

## 1 Introduction

The aim of this code library is to perform radiative transfer calculations for a range of applications, focusing however on the spectral synthesis of 3D hot-star winds. The main features consist of performing detailed radiative transfer calculations accounting for highly supersonic velocity fields and (almost) arbitrary 3D structures. We have developed several main programs to be used for different situations, summarized as follows:

**line3D:** A code module to calculate synthetic line profiles for a single star within a global *star-in-a-box* setup (see Sect. 3.1).

**BOSS-3D:** A code module to calculate synthetic line profiles for binary systems (see Sect. 3.3).

**cont3Dslab:** A code module to calculate the continuum radiation for a single star within a local *star-in-a-box* setup, that might be used for coupling in radiation-hydrodynamic simulations (see Sect. 3.2).

Each of these packages will be described in the corresponding sections in more detail.

## 1.1 Radiation hydrodynamics

ToDo

#### 1.2 Radiative transfer

To calculate the radiative transfer, we consider the time-independent equation of radiative transfer,

$$\boldsymbol{n}\nabla I_{\nu} = \eta_{\nu} - \chi_{\nu}I_{\nu} = \chi_{\nu}(S_{\nu} - I_{\nu}), \qquad (1)$$

with  $I_{\nu}$  the specific intensity,  $\eta_{\nu}$  the emissivity,  $\chi_{\nu}$  the opacity, and  $S_{\nu} = \eta_{\nu}/\chi_{\nu}$  the source function. Further, we define the angular moments of the specific intensity:

$$J_{\nu} = \frac{1}{4\pi} \int I_{\nu} d\Omega = \frac{c}{4\pi} E_{\nu}, \qquad (2)$$

$$\boldsymbol{H}_{\nu} = \frac{1}{4\pi} \int I_{\nu} \boldsymbol{n} d\Omega = \frac{1}{4\pi} \boldsymbol{F}_{\nu}, \tag{3}$$

$$K_{\nu} = \frac{1}{4\pi} \int \underbrace{nI_{\nu}n}_{\text{dvadic product}} d\Omega = \frac{c}{4\pi} P_{\nu},$$
 (4)

with  $J_{\nu}$  the mean intensity,  $H_{\nu}$  the Eddington flux, and  $K_{\nu}$  simply the second moment without a specific name. The mean intensity, Eddington flux, and second moment are trivially related to the radiation energy density  $E_{\nu}$ , the radiation flux  $F_{\nu}$ , and the radiation pressure tensor  $P_{\nu}$ .

For all code modules, we are typically considering the radiation quantities  $I_{\nu}$ ,  $J_{\nu}$ ,  $H_{\nu}$ ,  $K_{\nu}$ , and not the corresponding 'physical' quantities. In general there are three operating modes for calculating these quantities:

- (i) Since the source function in the equation of radiative transfer (EQRT) can in principle depend on the radiation field, Eq. (1) becomes an integro-differential equation. In this case, an iteration scheme is required (main code line3D/sc3d.eo and cont3Dslab/sc3d.eo for star-in-a-box and box-in-a-star simulations, respectively) based on a non-local accelerated Λ iteration (ALI).
- (ii) For known source functions and opacities (e.g., approximated in LTE or pre-calculated in step (i)), we can solve the radiative transfer in a *pz*-type geometry to obtain surface brightnesses or emergent flux profiles (main codes *line3D/modelspec.eo* and *line3D/spec.eo*).
- (iii) If we are dealing with binary systems, we rely purely on semi-analytical models thus far (e.g., opacities and source functions in LTE), and solve the radiative transfer in a *pz*-type of geometry (main codes *line3D/modelspec\_vbin.eo* and *line3D/spec\_vbin.eo*).

#### 1.3 Philosophy

All developed sub-programs are meant to be – at least in principle – a sort of stand-alone packages, that can be used completely independent of each other. Indeed, we consider the full radiative transfer problem as a three-step process:

1. Firstly, we need to create a discretized model of the physical state of the gas (i.e., density ρ, gas temperature T<sub>gas</sub>, velocity field v). In order that our radiative-transfer routines can communicate with such a model, we have developed a user interface to either transform input data from a given hydrodynamic simulation or to set up a semi-analytical model. This step is performed in the code model.eo with corresponding source code to be found in line3D/src\_model and cont3Dslab/src\_model.

- 2. Secondly, we need to calculate opacities and source functions. We can follow two branches here:
  - For resonance lines within a two-level-approximation and/or a two-component continuum source consisting of thermal and scattering terms, we can calculate continuum and line source functions consistently with the radiation field. This is performed by the code sc3d.eo. Since the iteration scheme is computationally very expensive, the source function will be calculated on a relatively low-resolution grid, and then interpolated back onto the original model within the code modelspec.eo. The corresponding source codes can be found in line3D/src\_sc3d, line3D/src\_modelspec, and cont3Dslab/src\_sc3d.
  - Alternatively, we can directly use the code *modelspec.eo* to calculate source functions and opacities from semi-analytical calculations (e.g., assuming LTE occupation numbers, see *line3D/src\_modelspec*).
- 3. Finally, the surface brightness or emergent flux profiles are calculated by solving the radiative transfer in a *pz*-type geometry using the code *spec.eo*, with input given from the previous step 2. The corresponding source code can be found in *line3D/src\_spec* and *cont3Dslab/src\_surfb*.

We emphasize that the binary version (extension \_vbin) consists only of steps (2) and (3), since we haven't implemented an ALI scheme for binary systems yet.

## 2 Installation

## 2.1 Requirements

The code requires the following packages:

**FORTRAN compiler:** Either the gfortran (version 8+) or ifort compiler is required. Depending which compiler is used, one needs to adapt the source code since the INQUIRE function works differently for both compilers: gfortran: inquire(file=trim(directory)//'/.', exist=my\_boolean) ifort: inquire(directory=trim(directory), exist=my\_boolean)

**HDF5:** The HDF5 library is required (version 1.10.5 or higher). This library needs to be compiled with the same compiler as used for the main programs (i.e., gfortran or ifort). See Sect. 2.4 on how to install HDF5 on your system

**PYTHON, IDL, GDL** The code comes with an IDL/GDL or PYTHON library for reading and plotting all output files.

## 2.2 Getting the code

You can get the code from github:

- Global star-in-a-box simulations: git clone https://github.com/levin-h/line3D
- Local box-in-a-star simulations: git clone https://github.com/levin-h/cont3D

## 2.3 Quick start

#### 2.4 Installing HDF5

I recommend to create a local build for your HDF5 libraries. To this end, please follow the following steps (here for version 1.10.6)

## **UNIX** systems

- 1. Download the package **hdf5-1.10.6.tar.gz**, unzip it, and change to the corresponding folder: tar -zxvf hdf5-1.10.6.tar.gz cd hdf5-1.10.6
- 2. Export some required environment variables:

```
gfortranifortexport FC=gfortranexport FC=ifortexport CC=gccexport CC=iccexport F9X=gfortranexport F9X=ifortexport CXX=g++export CXX=icpc
```

- 3. Configure your installation with a local path where you want to install the package (e.g., \$HOME/hdf5\_gfortran\_build): ./configure --prefix=\$HOME/hdf5\_gfortran\_build --enable-fortran (and if required --enable-cxx)
- 4. Installation

```
make (and watch for fatal errors)
make check (and verify that all tests return a 'pass')
make install
```

- 5. Include the library path in the Makefile of the main code (e.g., line3D/Makefile)
- 6. Update your .bashrc file to add the library path to your LD environment variable: HDF5\_PATH=\$HOME/hdf5\_gfortran\_build/hdf5\_lib/lib LD\_LIBRARY\_PATH=\$LD\_LIBRARY\_PATH:\$ADD\_LIB\_PATH export LD\_LIBRARY\_PATH

MAC On the MAC, the installation is essentially performed the same way. For the gfortran compiler, you might

- 1. Install the CommandLineTools
- 2. Install homebrew:

```
mkdir homebrew && curl -L https://github.com/Homebrew/brew/tarball/master | tar xz --strip 1 -C homebrew eval "$(homebrew/bin/brew shellenv)" brew update --force --quiet chmod -R go-w "$(brew --prefix)/share/zsh" export PATH=$HOME/homebrew/bin:$PATH
```

- 3. Install gfortran: brew install gcc
- 4. Install HDF5 as for UNIX systems. When updating the Makefile of the main code (e.g., line3D/Makefile), replace all \*.so libraries with the MAC \*.dylib extension.

## 2.5 Importing the plotting routines

All plotting routines are stored in the directory *line3D/plotFILES* or *cont3Dslab/plotFILES*, with corresponding libraries named *levpy*, *lib\_gdl* and *lib\_idl* for PYTHON, GDL, and IDL, respectively.

**GDL/IDL** The required libraries can be simply read in by typing in your IDL/GDL terminal: @idl\_lib/startup.pro @gdl\_lib/startup.pro

We emphasize that some of the routines might be outdated (for instance, variable names stored within the individual \*.h5 files might have changed). It should be straight forward though to adapt the corresponding reading routines (in *idl\_lib/proLEV/getall*).

**PYTHON** The following python packages are required: h5py, imageio, matplotlib, numpy, os, scipy, sys

## 3 Main code modules

#### **3.1** line3D

This folder contains code modules for running global (star-in-a-box) radiative transfer simulations. There are essentially four different code modules further described in the following. Each code module requires a specific input file organized by namelists.

**model.eo** Prepares the data to be used for the actual radiative transfer calculations, by transforming any input data or calculating semi-analytic models. You can simply add new models in src\_model. The corresponding namelist file is described in Table 1. Essentially, we set up the state of the gas in 1d, 2d, or 3d, described by the density  $\rho$ , the velocity field  $\boldsymbol{v}$ , the gas and radiation temperatures,  $T_{\rm gas}$  and  $T_{\rm rad}$ , the thermal velocities (becomes obsolete at some point),  $v_{\rm th}$ , and the thermalization parameter  $\epsilon_{\rm C}$ . Currently hardcoded, the model will be saved in inputFILES/modelXd.h5.

**sc3d.eo** Performs the radiative transfer for the continuum and/or line transition with certain opacity laws using an iterative ALI scheme. The corresponding namelist file is described in Table 1.

**modelspec.eo** Prepares the data to be used for the line-profile calculations of single stars. Here, we can read in the source functions and opacities calculated by the *sc3d.eo* program, or implement other semi-analytic models. The corresponding namelist file is described in Table 2.

**spec.eo** Calculates line profiles for a specific input file. The corresponding namelist files is described in Table 3.

#### 3.1.1 Program model.eo

All source files for this program are stored in the directory *src\_model*, and the corresponding namelist file is summarized in Table 1. Within this namelist, there are many input parameters that are actually not required. For consistency reasons and to avoid potential error sources, we decided to use the same namelist file also for the program *sc3d.eo*.

To register a new model, we recommend to follow the following steps:

- 1. In *src\_model/model.f90*, add a new model identifier as a case for the variable input\_mod. This identifier should be used also in the namelist file to call this particular model. The subroutine to create the model still needs to be developed by the user, e.g., *calc\_my\_model*, and needs to be called within the case of the new model identifier. Further, depending on the dimensionality of the new model, we need to save it as an h5 file by calling the (already existing) subroutines *output\_mod\*d*.
- In src\_model/model\*d.f90, we create our new subroutine calc\_my\_model. Depending on the dimension of the new model, different global variables have to be set (see src\_model/output\_model.f90 for more details). For a 3D model in spherical coordinates (r, Θ, Φ), we would require the following:

**nr\_modext** describes the number of radial grid points for our model.

**ntheta\_modext** describes the number of  $\Theta$  grid points for our model.

**nphi\_modext** describes the number of  $\Phi$  grid points for our model.

**r\_modext3d** describes the radial grid (array of length nr\_modext) in cgs.

**theta\_modext3d** describes the  $\Theta$  grid (array of length ntheta\_modext) from  $[0,\pi]$ .

**phi\_modext3d** describes the  $\Phi$  grid (array of length nphi\_modext) from  $[0, 2\pi]$ .

velr\_modext3d describes the radial velocity component (array of length nr\_modext, ntheta\_modext, nphi\_modext)
in cgs.

 $\begin{tabular}{ll} \textbf{velth\_modext3d} & describes the $\Theta$ velocity component (array of length nr\_modext, ntheta\_modext, nphi\_modext) \\ & in cgs. \end{tabular}$ 

**velphi\_modext3d** describes the  $\Phi$  velocity component (array of length nr\_modext, ntheta\_modext, nphi\_modext) in cgs.

rho\_modext3d describes the density (array of length nr\_modext, ntheta\_modext, nphi\_modext) in cgs.

 $t\_modext3d \ \ describes \ the \ gas \ temperature \ (array \ of \ length \ nr\_modext, \ ntheta\_modext, \ nphi\_modext) \ in \ cgs.$ 

trad\_modext3d describes the radiation temperature (array of length nr\_modext, ntheta\_modext, nphi\_modext)
in cgs. Often used only as a dummy array.

**vth\_modext3d** describes the thermal velocity (array of length nr\_modext, ntheta\_modext, nphi\_modext) in cgs. Often used only as a dummy array.

eps\_cont\_modext3d describes the thermalization parameter (array of length nr\_modext, ntheta\_modext, nphi\_modext).

To plot the resulting model, you can use the programs *plotFILES/model\*d.py* or *plotFILES/model\*d.pro* for PYTHON or IDL/GDL, respectively.

Table 1: Input namelist for the programs *model.eo* and *sc3d.eo*. Some of the inputs are meanwhile obsolete. We use one indat file for both programs to avoid inconsistency of the data used within *model.eo* and *sc3d.eo*. If only running the *model.eo* many of the parameters are not required and should be assigned with an arbitrary value.

Example	Data type	Description
&input_options		Options for the models
model_dir = 'inputFILES'	string	Directory of the model that will be read in
output_file = 'output_model00.h5'	string	All calculations stored in output_file (.h5 extension to be included)
input_mod = 12	integer	Identifier of the model to be calculated; only required for model.eo (where the hydro model is speci-
		fied)
$input\_mod\_dim = 3$	integer	Dimension of input model; input_mod_dim $\in [1,2,3]$
spatial_grid1d = 5	integer	Identifier to calculate a 1D radial grid from a beta-velocity law. Depending ong the option spa-
	2	tial_grid3d, this input is obsolete.

		spatial_grid1d=0 if equidistant radial grid is used (subroutine grid1d_r_equi)
		spatial_grid1d=1 if equidistant velocity grid is used (subroutine grid1d_vel_equi)
		spatial_grid1d=2 if equidistant tau_thomson grid is used (subroutine grid1d_tau_equi)
		spatial_grid1d=3 if equidistant log(tau_thomson) grid is used (subroutine grid1d_tau_log)
		spatial_grid1d=4 if combination is used (see subroutine grid1d_final for details)
		spatial_grid1d=5 if combination is used (see subroutine grid1d_final_2 for details)
	٠.	spatial_grid1d=6 if grid is calucalted equidistant in log-space (subroutine grid1d_r_log)
$spatial\_grid3d = 2$	integer	Identifier to calculate the 3D Cartesian grid.
		spatial_grid3d=0 if 3d grid is calculated from 1d grid with equidistant core points spatial_grid3d=1 if 3d grid is calculated from a mean-value approach (minimizing distance of subse-
		quent coordinates from 1d-grid)
		spatial_grid3d=2 if 3d grid is calculated from a mean-value approach (minimizing distance of subse-
		quent coordinates from original input-grid)
		spatial_grid3d=3 if 3d grid is calculated completely equidistant
		spatial_grid3d=4 if 3d grid is calculated from a 1d radial grid and setting up angular grid equidistantly
		spatial_grid3d=5 if 3d grid is calculated from a 3d spherical grid (optimized)
$opt\_opac = 0$	integer	Identifier to decide on the continuum opacity model
•		opt_opac=0 if Thomson opacities
		opt_opac=1 if OPAL opacities
$opt\_opal = 0$	integer	Identifier to decide on the line opacity model
		opt_opal=0 if line-strength parameter
		opt_opal=1 if Hamann (1980) parameterization
$opt\_angint\_method = 9$	integer	Identifier to decide on the angular-integration technique to be used
		opt_angint_method=0 if angular integration is used with trapezoidal rule (nodes equidistant in $\theta$ and
		$\phi$ )
		opt_angint_method=1 if angular integration is used with trapezoidal rule (nodes from Lobel &
		Blomme (2008))
		opt_angint_method=2 if angular integration is used with simpsons rule (nodes equidistant in $\theta$ and $\phi$ ,
		note: $\mu$ -grid and $\phi$ -grid will be made equidistant for three subsequent points)
		opt_angint_method=3 if angular integration is used with simpson rule corrected for the error from a
		grid with half resolution (also known as boole's rule)
		opt_angint_method=4 if angular integration is used with cubic splines (catmull-rom-spline, nodes
		equidistant in $\theta$ and $\phi$ )
		opt_angint_method=5 if angular integration is used with gauss-legendre-integration (for each octant) opt_angint_method=6 if angular integration is used with gauss-chebyshev-integration (for each oc-
		tant)
		opt_angint_method=7 if angular integration is used with triangulation (linear integrals)
		opt_angint_method=8 if angular integration is used with triangulation ('pseudo'-gauss integrals per
		triangle)
		opt_angint_method=9 if angular integration is used with lebedev interpolation (optimized nodes on
		the sphere)
$opt\_method = 1$	integer	Identifier to decide on the radiative-transfer solution method
. –	C	opt_method=0 if finite volume method shall be used
		opt_method=1 if linear short characteristics method shall be used
		opt_method=2 if quadratic bezier short characteristics method shall be used
$opt\_sol2d = f$	logical	Logical to decide whether 2D solution scheme shall be applied
$opt\_ltec = 0$	integer	Identifier to decide on the continuum wavelength/frequency model
	ınteger	
	integer	opt_ltec = 0 if single continuum frequency
	integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the
	integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium
		opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ).
opt_incl_cont = t	logical	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not)
opt_start_cont = t	logical logical	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps)
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<pre>opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f</pre>	logical logical logical logical logical	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not)
<pre>opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t</pre>	logical logical logical logical logical logical	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps)
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<pre>opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t opt_ng_line = t</pre>	logical logical logical logical logical logical logical	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration
<pre>opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t opt_ng_line = t opt_ait_line = f</pre>	logical logical logical logical logical logical logical logical	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration
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<pre>opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t opt_ng_line = t opt_ait_line = f</pre>	logical logical logical logical logical logical logical logical	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 1 if diagonal approximate $\Lambda$ operator (7 elements)
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opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t opt_ng_line = t opt_ait_line = f opt_ait_line = f	logical logical logical logical logical logical logical integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if hie shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_cont = 3 if nearest-neighbour approximate $\Lambda$ operator (27 elements) Identifier to define the approximate $\Lambda$ -operator for line iteration opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator
opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t opt_ng_line = t opt_ait_line = f opt_ait_line = f	logical logical logical logical logical logical logical integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_cont = 3 if nearest-neighbour approximate $\Lambda$ operator (27 elements) Identifier to define the approximate $\Lambda$ -operator for line iteration opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator (7 elements)
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<pre>opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t opt_ng_line = t opt_ait_line = f opt_alo_cont = 3</pre> opt_alo_line = 3	logical logical logical logical logical logical integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_cont = 3 if nearest-neighbour approximate $\Lambda$ operator (27 elements) Identifier to define the approximate $\Lambda$ -operator for line iteration opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (27 elements) Set to true (false) if gravity darkening by von Zeipel (1924) shall be included (or not)
<pre>opt_start_cont = t opt_ng_cont = t opt_ait_cont = f opt_incl_line = f opt_start_line = t opt_ng_line = t opt_ait_line = f opt_alo_cont = 3  opt_alo_line = 3</pre>	logical logical logical logical logical logical logical integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 1 if diagonal approximate $\Lambda$ operator opt_alo_cont = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator opt_alo_line = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) Set to true (false) if gravity darkening by von Zeipel (1924) shall be included (or not)
opt_start_cont = t  opt_ng_cont = t  opt_ait_cont = f  opt_incl_line = f  opt_start_line = t  opt_ait_line = f  opt_alo_cont = 3  opt_alo_line = 3  opt_incl_gdark = f  opt_incl_sdist = f  &input_mod_ld	logical logical logical logical logical logical integer  integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 1 if diagonal approximate $\Lambda$ operator opt_alo_cont = 2 if direct-neighbour approximate $\Lambda$ operator (27 elements) Identifier to define the approximate $\Lambda$ -operator for line iteration opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator (7 elements) opt_alo_line = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) Set to true (false) if gravity darkening by von Zeipel (1924) shall be included (or not) Input parameters of the considered star (some not required anymore)
opt_start_cont = t  opt_ng_cont = t  opt_ait_cont = f  opt_incl_line = f  opt_start_line = t  opt_ait_line = f  opt_alo_cont = 3  opt_alo_cont = 3  opt_alo_line = 3	logical logical logical logical logical logical integer  integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if Continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 1 if diagonal approximate $\Lambda$ operator (7 elements) opt_alo_cont = 2 if direct-neighbour approximate $\Lambda$ operator (27 elements) Identifier to define the approximate $\Lambda$ -operator for line iteration opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator (7 elements) opt_alo_line = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) Set to true (false) if gravity darkening by von Zeipel (1924) shall be included (or not) Set to true (false) if surface distortion due to rotation shall be included (or not) Effective temperature of the star in [K]
opt_start_cont = t  opt_ng_cont = t  opt_ait_cont = f  opt_incl_line = f  opt_start_line = t  opt_ait_line = f  opt_alo_cont = 3  opt_alo_line = 3  opt_incl_gdark = f  opt_incl_sdist = f  &input_mod_ld	logical logical logical logical logical logical integer  integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Ng-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 1 if diagonal approximate $\Lambda$ operator opt_alo_cont = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_lone = 0 if classical $\Lambda$ iteration opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator opt_alo_line = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) Set to true (false) if surface distortion due to rotation shall be included (or not) Set to true (false) if surface distortion due to rotation shall be included (or not) Effective temperature of the star (used as the inner boundary condition for the specific intensity) in
opt_start_cont = t  opt_ng_cont = t  opt_ait_cont = f  opt_incl_line = f  opt_start_line = t  opt_ait_line = f  opt_alo_cont = 3  opt_alo_cont = 3  opt_alo_line = 3	logical logical logical logical logical logical integer  integer	opt_ltec = 0 if single continuum frequency opt_ltec = 1 if grey approximation for continuum (frequency integrated). If this option is set, the temperature will be updated after the radiation-transfer calculations assuming radiative equilibrium (i.e., from $J = S = B = \sigma_B/\pi T^4$ ). Set to true (false) if continuum shall be included (or not) Set to true (false) if Continuum iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for continuum iteration shall be included or not Set to true (false) if Aitkens-extrapolation for continuum iteration shall be included or not Set to true (false) if line shall be included (or not) Set to true (false) if line iteration shall start from the beginning (or from intermediate steps) Set to true (false) if Ng-extrapolation for line iteration shall be included or not Set to true (false) if Aitkens-extrapolation for line iteration shall be included or not Identifier to define the approximate $\Lambda$ -operator for continuum iteration opt_alo_cont = 0 if classical $\Lambda$ iteration opt_alo_cont = 1 if diagonal approximate $\Lambda$ operator (7 elements) opt_alo_cont = 2 if direct-neighbour approximate $\Lambda$ operator (27 elements) Identifier to define the approximate $\Lambda$ -operator for line iteration opt_alo_line = 0 if classical $\Lambda$ iteration opt_alo_line = 1 if diagonal approximate $\Lambda$ operator (7 elements) opt_alo_line = 2 if direct-neighbour approximate $\Lambda$ operator (7 elements) opt_alo_line = 3 if nearest-neighbour approximate $\Lambda$ operator (7 elements) Set to true (false) if gravity darkening by von Zeipel (1924) shall be included (or not) Set to true (false) if surface distortion due to rotation shall be included (or not) Effective temperature of the star in [K]

rstar = 8.d0	float	$R_*$ in $R_{\odot}$
lstar = 1.d6	float	$L_*$ in $L_\odot$
rmax = 12.d0	float	Maximum radius of the computational domain in $[R_*]$ along each $x, y, z$ axis
tmin = .8d0	float	Minimum temperature in the wind in $[T_{rad}]$
xmloss = 5.d-6	float	mass-loss rate $\dot{M}$ in $M_{\odot}$ yr <sup>-1</sup> ; only required for 1D benchmarking
vmin = 1.d1	float	minimum velocity of $\beta$ -velocity law $v_{min}$ in km s <sup>-1</sup> ; only required for 1D benchmarking
vmax = 2.d3	float	terminal velocity of $\beta$ -velocity law $v_{\infty}$ in kms <sup>-1</sup> ; only required for 1D benchmarking
vmicro = 1.d2	float	micro-turbulent velocity for the line-profile function $v_{\text{turb}}$ in [km s <sup>-1</sup> ]
	float	
vth_fiducial= 1.d2		fiducial thermal velocity $v_{\text{th}}^*$ in $[\text{km s}^{-1}]$ rotational velocity $v_{\text{rot}}$ in $[\text{km s}^{-1}]$
vrot = 0.d0 $beta = 1.d0$	float	
	float	$\beta$ parameter for $\beta$ -velocity law; only required for 1D benchmarking models
yhe = .1d0 hei = 2.d0	float	Helium abundance by number, Y <sub>He</sub>
xnue0 = 1.93798d15	float float	Helium ionization fraction (number of free electrons per Helium-atom)
na = 12		Frequency of the line transition mass number A for the line transition
	integer	
&input_infreg	floot	Input parameters to define the computational domain (information region)
rmin = 1.d0 $rlim = 13.2d0$	float	Minimum radius of the computational domain in $R_*$
rlim = 13.2d0	float	Maximum radius of the computational domain in $R_*$
&input_cont	α .	Parameters for the continuum transport
eps_cont = 0.d0	float	Thermalization parameter $\epsilon_{\mathrm{C}}$
kcont = 1.d0	float	$k_C$ parameter (linear scaling factor for the continuum opacity)
&input_line	g4	Parameters of the line transport
eps_line = 0.d0	float	Line-scattering parameter $\epsilon_{ m L}$
kline = 1.d0	float	line-strength parameter $k_{\rm L}$
kappa0 = 1.d-1	float	Hamann (1980) parameterization
alpha = 0.5d0	float	Hamann (1980) parameterization
&dimensions_1d		Dimension parameters to set up 1D radial grid
n1d = 17	integer	number of radial grid points (used to distribute z-axis in $[R_{\min}, R_{\max}]$ )
$n1d_{-}t = 81$	integer	number of 1D grid points to set up equidistant $\tau$ -grid
$n1d_r = 22$	integer	number of 1D grid points to set up equidsitant $v_r$ -grid
$\frac{\text{delv} = 0.33d0}{\frac{1}{10000000000000000000000000000000$	float	Preferred velocity steps $\Delta v_r$ in $v_{th}^*$
&dimensions_3d	• .	Dimension parameters to set up the 3D grid
ncx=19	integer	Preferred number of core-points for x-axis
ncy=19	integer	Preferred number of core-points for y-axis
ncz=19	integer	Preferred number of core-points for z-axis
delx_max=.7d0	float	Maximum allowed $\Delta x$ in $R_*$
dely_max=.7d0	float	Maximum allowed $\Delta y$ in $R_*$
delz_max=.7d0	float	Maximum allowed $\Delta z$ in $R_*$
&dimensions_freq	g .	Dimension parameters to set up the frequency grid
deltax = 0.333d0	float	$\Delta x_{\rm obs}$ steps
$xcmf_max = 3.d0$	float	Maximum frequency width of the line-profile function, $x_{cmf}^{(max)}$
&dimensions_angles		Dimension parameters to set up the angular grid
$n_{theta} = 11$	integer	Number of $\theta$ angles in first octant; $\phi$ angles are calculated based on that
&benchmark		Parameters for setting up a benchmark
$benchmark\_mod = 0$	integer	Identifier to define the benchmark model (set to 0 if no benchmark shall be performed)
im_source = 3	integer	see benchmark subroutines
im_opacity = 2	integer	see benchmark subroutines
$im_vel = 0$	integer	see benchmark subroutines
$tau_min = 0.d0$	float	see benchmark subroutines
$tau_max = 5.d0$	float	see benchmark subroutines
$source\_min = 0.1d0$	float	see benchmark subroutines
$source_max = 1.d-6$	float	see benchmark subroutines
$n_y = 0.d0$	float	see benchmark subroutines
$n_z = 0.707107d0$	float	see benchmark subroutines

#### 3.1.2 Program sc3d.eo

This program solves the non-linear coupling of the radiative transfer equation with the source function of the form:

$$S_{C} = (1 - \epsilon_{C})J_{\nu} + \epsilon_{C}B_{\nu} \tag{5}$$

$$S_{L} = (1 - \epsilon_{L})\bar{J} + \epsilon_{L}B_{\nu_{0}}, \tag{6}$$

i.e., for a continuum consisting of thermal and scattering terms, and for a resonance-line transition approximated as a two-level atom. To this end, we are discretizing the equation of radiative transfer in Cartesian coordinates, and rely on the accelerated  $\Lambda$ -iteration (ALI) using non-local approximate  $\Lambda$  operators (ALO). The corresponding source files can be found in  $src\_sc3d$ .

There are various different methods for solving the radiative transfer equation (e.g., via the finite-volume method or the short-characteristics method), as well as for performing the source-function updates (using different ALO's). All available options required for the input namelist are summarized in Table 1. As output and depending on the chosen options, the \*.h5 file generated by sc3d.eo provides among other data:

**scont3d** The continuum source function  $S_C$  in cgs (3d array with dimensions (nx,ny,nz)).

**mint3d** The mean intensity  $J_{\nu}$  in cgs (3d array with dimensions (nx,ny,nz)).

**fcontx3d, fconty3d, fcontz3d** The Eddington flux components in Cartesian coordinates,  $\mathbf{H}_v = (H_x, H_y, H_z)$ , in cgs (3d array with dimensions (nx,ny,nz)).

**kcontxx3d, kcontyy3d, kcontxz3d, kcontxx3d, kcontxz3d, kcontyz3d** The tensor components of the  $K_{\nu}$ -tensor (3d arrays with dimensions (nx,ny,nz)). We emphasize that this is a symmetric tensor, and only six components need to be saved to deduce the complete tensor.

**mintbar3d** The frequency integrated and profile weighted mean intensity,  $\bar{J}$  in cgs (3d array with dimensions (nx,ny,nz)).

**sline3d** The line source function,  $S_L$  in cgs (3d array with dimensions (nx,ny,nz)).

To plot the resulting model, we provide the PYTHON and GDL/IDL programs *plotFILES/plot\_sc3d.py* and *plot-FILES/plot\_sc3d.pro*.

#### 3.1.3 Program modelspec.eo

This program prepares data to be used for calculating spectral features and/or surface brightnesses of our simulations. We can either calculate a line profile from the output (i.e., source functions) of the *sc3d.eo* program, or create a completely new semi-analytic model. All source files can be found in the *src\_modelspec*/ directory, with the available namelist options summarized in Table 2.

To create a new model here, we essentially follow the same philosophy as for the *model.eo* program, and recommend the following two steps:

- In *src\_modelspec/modelspec.f90*, we can add a new model identifier as a case for the namelist variable in-uput\_mod. As before, we can then create and call a new subroutine describing our model.
- In *src\_modelspec/modelspec.f90*, we also create the new subroutine, e.g., *subroutine my\_model*. Within this subroutine (or in the input namelist), we need to specify the following global variables (here for a standard 3D model in spherical coordinates).

nr Number of radial grid points.

**ntheta** Number of  $\Theta$  grid points.

**nphi** Number of  $\Phi$  grid points.

**r** The radial grid in  $R_*$ .

**theta** The  $\Theta$  grid in the range  $[0,\pi]$ .

**phi** The  $\phi$  grid in the range  $[0, 2\pi]$ .

sline3d The line source function in cgs (3d array with dimensions (nr,ntheta,nphi)).

scont3d The continuum source function in cgs (3d array with dimensions (nr,ntheta,nphi)).

t3d The gas temperature in cgs (3d array with dimensions (nr,ntheta,nphi)).

**opac3d** The continuum opacity in  $[1/R_*]$  (3d array with dimensions (nr,ntheta,nphi)).

**oplbar3d** The frequency integrated line opacity in  $[1/sR_*]$  (3d array with dimensions (nr,ntheta,nphi)).

velx3d The x-component of the velocity field in cgs (3d array with dimensions (nr,ntheta,nphi)).

vely3d The y-component of the velocity field in cgs (3d array with dimensions (nr,ntheta,nphi)).

velz3d The z-component of the velocity field in cgs (3d array with dimensions (nr,ntheta,nphi)).

xic1, xic2 The anchor for the inner boundary condition of the specific intensity, which should follow the form for core rays:

$$I_{\nu} = xic1 \cdot q_1 - xic2 \cdot q_2, \tag{7}$$

where  $q_1$  and  $q_2$  are scaling factors to be calculated during the formal solution (e.g.,  $q_1$  can be set to account for gravity darkening). A reasonable choice, for instance might be:

$$xic1 = B_{\nu}(T_{\text{eff}})$$
  $xic2 = \frac{dB\nu}{\chi_{\nu}dz}$ . (8)

Again, we can display the resulting model by using the programs plotFILES/modelspec3d.py or plotFILES/modelspec3d.pro.

Table 2: Input namelist for the program *modelspec.eo* 

Example	Data type	Description
&input_options		Main options
input_file =	string	Name of the input file generated by sc3d.eo, if source funtions and opacities are to be read in from the
'./outputFILES/output_model00.h5'		solution of sc3d.eo
input_file2 =	string	Name of the input model file generated by model.eo. Depending on the input_mod options, all opac-
'./inputFILES/model3d.h5'		ities and source functions are either interpolated from the sc3d.eo output onto this grid, or calculated
		from a semi-analytical model. This procedure allows us to use a low-resolution grid for the compu-
		tationally challenging ALI iteration, while still using a high-resolution grid of the wind's density and
		velocity structure.
output_file =	string	Output file
'./outputFILES/modspec_model00.h5	5'	
input_mod = 19	integer	Identifier for the model to be calculated (see in ./src_modelspec/modelspec.f90). There are a few
-	_	standard options to communicate with the output from the program sc3d.eo, such as:
		input_mod=11 3d model: standard ouput from sc3c.eo (3d cartesian model)
		input_mod=12 3d model: standard ouput from sc3c.eo (3d cartesian model) interpolated onto the
		spherical grid from the model.eo output
&input_model		Parameters of the input model
teff = 258390.7d0	float	Effective temperature of the star. Only required to get the correct photospheric line profile later on.
trad = 258390.7d0	float	Radiation temperature of the star. Only used to set the inner boundary condition for the specific
		intensity.
xlogg = 3.6d0	float	$\log g$ of the star. Only used to get the correct photospheric line profile later on.
rstar = 1.d0	float	$R_*$ in $R_{\odot}$
rmax = 11.d0	float	$R_{\text{max}}$ in $R_*$ , used to define the computational domain
tmin = 1.d0	float	Minimum temperature of the wind in $[T_{\text{eff}}]$ . Only used for very specific test routines.
xmloss = 1.d-6	float	Mass-loss rate $\dot{M}$ in $[M_{\odot} yr^{-1}]$ . Only used for very specific test routines.
vmin = 10.d0	float	Minimum velocity $v_{\min}$ of a $\beta$ -velocity law in [km s <sup>-1</sup> ]. Only used for very specific test routines
vmax = 4.d3	float	Terminal velocity $v_{\infty}$ of a $\beta$ -velocity law in [km s <sup>-1</sup> ]. If not overwritten within the specific model
		routines, this sets also the range of velocities/frequencies for which the line-profiles are calcu-
		lated
beta = 1.d0	float	$\beta$ parameter of a $\beta$ -velocity law in [km s <sup>-1</sup> ]. Only used for very specific test routines
vmicro = 1.0d2	float	Microturbulent velocity $v_{\text{turb}}$ in [km s <sup>-1</sup> ].
vth_fiducial=1.d2	float	Fiducial thermal velocity to be used in [km s <sup>-1</sup> ].
yhe = 0.1d0	float	Helium number abundance, $Y_{\text{He}} = n_{\text{He}}/n_{\text{H}}$ (e.g., $Y_{\text{He}} = 12.25$ corresponds to mass-fraction 0.98).
hei = 2.d0	float	Number of free electrons per helium atom
&input_line		Line parameters
iline = 0	integer	Identifier for the line (as defined in src/mod_iline.f90) to get all line data $(v_0, g_l, g_u, \text{etc})$
		iline=0 - read atomic charge Z, element i, lower level l and upper level u from file 'in_linelist.dat'
		iline=1 - $H_{\alpha}$
		iline=2 - $H_{\beta}$
		iline=10 - CIV resonance line
		iline=11 - C III 5696 line
$eps_line = 0.d0$	float	Line scattering parameter $\epsilon_L$ . Only used for specific test routines (Sobolev solution)
kline = 1.d0	float	Line-strength parameter or arbitrary scaling factor to increase/decrease the line opacity
kappa0 = 1.d0	float	Hamann (1980) parameterization
alpha = 0.d0	float	Hamann (1980) parameterization

## 3.1.4 Program spec.eo

This program calculates synthetic line profiles and surface brightnesses for a given model obtained by the program *modelspec.eo*. To this end, we rely on a cylindric coordinate system  $(p, \zeta, z)$  (see also Hennicker et al. (2021)). When calculating surface brightnesses the output will be stored as \*.h5 file giving:

**p** The array of impact parameters.

zeta The array of angles of the cylindrical coordinate system

**iem\_surface** The (total) emergent intensity at each  $p, \zeta$  in cgs.

**iemi\_surface** The emission part of the total intensity at each  $p, \zeta$  in cgs.

**iabs\_surface** The absorption part of the total intensity at each  $p, \zeta$  in cgs.

**icont\_surface** The continuum intensity only (if there was no line) at each  $p, \zeta$  in cgs.

When calculating emergent flux profiles, the output will be stored as ASCII files in *FLUXEM\_\*.dat*. The output is organized in columns giving:

**xobs** The frequency shift from line center in units of the fiducial velocity  $v_{th}^*$ .

**flux\_tot** The total emergent flux-like (or rather luminosity-like) quantity at this frequency. Following, e.g., Hennicker et al. (2020, Sect. 3.7), the flux is given by:

$$F_{\nu} = \frac{1}{d^2} \underbrace{\int_0^{2\pi} \int_0^{R_{\text{max}}} I_{\nu}(p, \zeta, z = R_{\text{max}}) \, p \mathrm{d}p \mathrm{d}\zeta}_{=:\text{flux\_tot}}. \tag{9}$$

Since we have been integrating over the impact parameter p (which internally is measured in  $R_*$ ), we can translate the quantity flux\_tot to a luminosity in cgs:

$$L_{\nu} = \text{flux\_tot} \cdot R_*^2 \cdot 4\pi. \tag{10}$$

The namelist options for the program spec.eo are summarized in Table 3.

Table 3: Input namelist for the program spec.eo

Data type	Description
	Main options
integer	Type of the input model
	$input\_mod = 0 - 1D \mod e$ on radial grid
	$input\_mod = 1 - 3D \mod on Cartesian grid$
	$input\_mod = 2 - 3D \mod on spherical grid$
string	Name of the input file generated by <i>modelspec.eo</i>
5'	
string	Output directory
C	
integer	Identifier for defining the photospheric line profile
	opt_photprof = 0 - no photospheric line profile (flat illumination)
	opt_photprof = 1 - from A. Herrero files
	opt_photprof = 2 - from Kurucz (not active at the moment)
	opt_photprof = 3 - from own FASTWIND compilation (only active in the binary version at the mo-
	ment)
	opt_photprof = 4 - from Coelho et al. (2005) (only active in the binary version at the moment)
	opt_photprof = 5 - from Coelho (2014) (only active in the binary version at the moment)
logical	Logical to decide whether observer's direction shall be read in or calculated.
	opt_obsdir_read = t - read in angles $\alpha \in [0,180]$ (measured from the z-axis, inclination) and $\gamma \in$
	[0,360] (measured from the x-axis, phase) from files in_alpha.dat and in_gamma.dat
	opt_obsdir_read = f - Equidistant $\alpha$ , $\gamma$ grid will be calculated based on input options nalpha and
	ngamma.
logical	Logical to decide if surface brightness shall be calculated instead of emergent flux profiles.
logical	Logical to decide if the propagation of intensity along a 2D slice trough the computational domain
_	shall be calculated instead of emergent flux profiles
logical	Logical to decide if von Zeipel (1924) gravity darkening shall be included
logical	Logical to decide if surface distortion shall be accounted for
integer	Number of $\alpha$ angles to define the directions to the observer
integter	Number of $\gamma$ angles to define the directions to the observer
	Input parameters for the model
float	Surface rotation of the star in $[km s^{-1}]$ (at the equator).
float	Fiducial thermal velocity $v_{th}^*$ in [km s <sup>-1</sup> ].
float	Microturbulent velocity $v_{\text{turb}}$ in[km s <sup>-1</sup> ].
float	Minimum radius of the computational domain (as used for <i>modelspec.eo</i> ).
float	Maximum radius of the computational domain (typically a bit smaller than used for modelspec.eo to
	avoid extrapolation errors/interpolations to zero).
	Input parameters for surface brightness calculations and calculating intensities along a 2d slice. Will
	be used only if either opt_surface or opt_int2d is set to true
integer	Number of surface brightnesses to be calculated.
float	The $\alpha$ angles towards the observer (number of elements needs to be equal to nsurfb).
float	The $\gamma$ angles towards the observer (number of elements needs to be equal to nsurfb).
	•
float	The shift from line center in units of $v_{th}^*$ (number of elements needs to be equal to nsurfb)
	ш
	For this example, two surface brightnesses will be calculated with directions and frequencies taken
	For this example, two surface originalesses will be calculated with directions and frequencies taken
	integer  string 5's string integer  logical logical logical integer integter  float

## 3.2 boss3D

The BOSS-3D package (also within the *line3D* folder with corresponding programs *line3D/modelspec\_vbin.eo* and *line3D/spec\_vbin.eo*, the extension *vbin* abbreviating 'version binary') contains modules for running global (star-in-abox) radiative transfer simulations of binary systems. There are two different code modules further described in the following. Each code module requires a specific input file organized by namelists.

**modelspec\_vbin.eo** Prepares the data to be used for the line-profile calculations of binary systems. Here, we define the model in terms of density, temperature and velocity fields, as well as opacities and source functions. The corresponding namelist file is described in Table 4.

**spec\_vbin.eo** Calculates line profiles for a specific input file. The corresponding namelist file is described in Table 5.

#### 3.2.1 The namelist files

The namelist options for the programs *modelspec\_vbin.eo*, and *spec\_vbin.eo* are shown in Tables 4 and 5, respectively.

Table 4: Input namelist for the program *modelspec\_vbin.eo*, i.e., for the binary version.

Example	Data type	Description
&input_options	71	Main options
input_file = "	string	Name of the input file generated by sc3d.eo (not used yet in binary version)
input_file2 = "	string	Name of the input model file generated by <i>model.eo</i> (not used yet in binary version)
output_file =	string	Output file
'./outputFILES/modspec_mode	el00.h5'	
$input\_mod = 9$	integer	Identifier for the model to be calculated (see in ./src_modelspec_vbin/modelspec.f90).
&input_model1		Parameters of the input model for the primary object.
rstar1 = 1.d0	float	$R_*^{(1)}$ of the primary object in $R_{\odot}$ , defining the length scale of the coordinate system of the primary.
rmin1 = 1.d0	float	Minimum radius defining the computational domain of the primary object, $R_{\min}$ in $R_*^{(1)}$
rmax1 = 10.d0	float	Maximum radius defining the computational domain of the primary object, $R_{\text{max}}$ in $R_*^{(1)}$
teff1 = 6.d3	float	Effective temperature of the primary object. Only required to get the correct photospheric line profile
		later on.
trad1 = 6.d3	float	Radiation temperature of the primary object. Only used to set the inner boundary condition for the
		specific intensity.
logg1 = 1.d0	float	$\log g$ of the primary object. Only used to get the correct photospheric line profile later on.
yhe1 = 0.1d0	float	Helium number abundance, $Y_{\text{He}} = n_{\text{He}}/n_{\text{H}}$ , of the primary object.
fehe1 = -1.d0	float	Fe/He abundance of primary object. Only required for Coelho et al. (2005) and Coelho (2014) photo-
		spheric line profiles.
aenh1 = 0.d0	float	$\alpha$ -element enhancement of primary object. Only required for Coelho et al. (2005) and Coelho (2014)
		photospheric line profiles.
vrot1 = 10.d0	float	Surface rotation of the primary object in [km s <sup>-1</sup> ] (at its equator).
vmicro1 = 1.0d2	float	Microturbulent velocity of the primary object $v_{\text{turb}}$ in [kms <sup>-1</sup> ].
p_object01 =	float	x,y,z position of the primary object within the global (center-of-mass) coordinate system in units
0.d0, 3.d0, 0.d0	~	[unit_length] (see &input_units)
v_object01 =	float	$v_x, v_y, v_z$ components of the orbit of the primary object within the global center-of-mass coordinate
-10.d0, 0.d0, 0.d0	g ,	system in [kms <sup>-1</sup> ].
ex01 = 1.d0, 0.d0, 0.d0	float	Orientation of the $e_x$ basis vector of the primary object within the global center-of-mass coordinate
21/01 - 0 40 1 40 0 40	float	system.  Orientation of the a hosis vector of the minory chiest within the clobal center of mass coordinate.
ey01 = 0.d0, 1.d0, 0.d0	noat	Orientation of the $e_y$ basis vector of the primary object within the global center-of-mass coordinate
ez01 = 0.d0, 0.d0, 1.d0	float	system. Orientation of the $e_z$ basis vector of the primary object within the global center-of-mass coordinate
ezo1 = 0.do, 0.do, 1.do	noat	System.
rot_axis01 =	float	Orientation of the rotation axis of the primary object within the global center-of-mass coordinate
0.d0, 0.d0, 1.d0	nout	system (still to be implemented).
&input_model2		Parameters of the input model for the secondary object. Same as for primary object but interchaning
compat_model2		the variable name index 1 with 2.
rstar2 = 3.d0	float	$R_*^{(2)}$ of the secondary object in $R_{\odot}$ , defining the length scale of the coordinate system of the secondary.
rmin2 = 3.d0 $rmin2 = 1.d0$	float	Minimum radius defining the computational domain of the secondary object, $R_{\text{min}}$ in $R_*^{(2)}$
rmax2 = 100.d0	float	Maximum radius defining the computational domain of the secondary object, $R_{\text{max}}$ in $R_*^{(2)}$
teff2 = 10.d3	float	Effective temperature of the secondary object. Only required to get the correct photospheric line
teli2 = 10.d3	noat	profile later on.
trad2 = 10.d3	float	Radiation temperature of the secondary object. Only used to set the inner boundary condition for the
11442 = 10.43	nout	specific intensity.
$\log g2 = 3.d0$	float	$\log g$ of the secondary object. Only used to get the correct photospheric line profile later on.
yhe2 = 0.1d0	float	Helium number abundance, $Y_{\text{He}} = n_{\text{He}}/n_{\text{H}}$ , of the secondary object.
fehe2 = -1.d0	float	Fe/He abundance of secondary object. Only required for Coelho et al. (2005) and Coelho (2014)
		photospheric line profiles.
aenh2 = 0.d0	float	$\alpha$ -element enhancement of secondary object. Only required for Coelho et al. (2005) and Coelho (2014)
		photospheric line profiles.
vrot2 = 100.d0	float	Surface rotation of the secondary object in [km s <sup>-1</sup> ] (at its equator).
vmicro2 = 1.0d1	float	Microturbulent velocity of the secondary object $v_{\text{turb}}$ in [km s <sup>-1</sup> ].
p_object02 =	float	x,y,z position of the secondary object within the global (center-of-mass) coordinate system in units
0.d0, -2.d0, 0.d0		[unit_length] (see &input_units)
v_object02 =	float	$v_x, v_y, v_z$ components of the orbit of the secondary object within the global center-of-mass coordinate
6.d0, 0.d0, 0.d0		system in $[km s^{-1}]$ .
ex02 = 1.d0, 0.d0, 0.d0	float	Orientation of the $e_x$ basis vector of the secondary object within the global center-of-mass coordinate
		system.
ey02 = 0.d0, 1.d0, 0.d0	float	Orientation of the $e_y$ basis vector of the secondary object within the global center-of-mass coordinate
0.0000000000000000000000000000000000000	~	system.
ez02 = 0.d0, 0.d0, 1.d0	float	Orientation of the $e_z$ basis vector of the secondary object within the global center-of-mass coordinate
		cyctem

system.

rot_axis01 =	float	Orientation of the rotation axis of the secondary object within the global center-of-mass coordinate
0.d0, 0.d0, 1.d0		system (still to be implemented).
&input_line		Line parameters
iline = 0	integer	Identifier for the line (as defined in src/mod_iline.f90) to get all line data ( $v_0, g_1, g_u$ , etc)
		iline=0 - read atomic charge $Z$ , element $i$ , lower level $l$ and upper level $u$ from file 'in_linelist.dat'
		iline=1 - $H_{\alpha}$
		iline=2 - $H_{\beta}$
		iline=10 - C IV resonance line
		iline=11 - C III 5696 line
$eps\_line = 0.d0$	float	Line scattering parameter $\epsilon_L$ . Only used for specific test routines (Sobolev solution)
kline = 1.d0	float	Line strength parameter
&input_units		Units of the simulation
$unit_length = 1.d0$	float	Length scale of the global coordinate system in $[R_{\odot}]$
$vth_fiducial = 1.0d2$	float	Fiducial thermal velocity $v_{th}^*$ .

Table 5: Input namelist for the program *spec\_vbin.eo*, i.e., for the binary version

Example	Data type	Description
&input_options		Main options
$input\_mod = 2$	integer	Type of the input model
		$input\_mod = 2 - 3D \mod on spherical grid$
input_file =	string	Name of the input file generated by <i>modelspec_vbin.eo</i>
'./outputFILES/modspec_model(	00.h5'	
output_dir =	string	Output directory
'./outputFILES'		
opt_photprof1 = 5	integer	Identifier for defining the photospheric line profile of the primary object.
opt_photprof2 = 0	integer	Identifier for defining the photospheric line profile of the secondary object.
1 –1 1	C	opt_photprof = 0 - no photospheric line profile (flat illumination)
		opt_photprof = 1 - from A. Herrero files
		opt_photprof = 2 - from Kurucz (not active at the moment)
		opt_photprof = 3 - from own FASTWIND compilation
		opt_photprof = 4 - from Coelho et al. (2005)
		opt_photprof = 5 - from Coelho (2014)
opt_obsdir_read = t	logical	Logical to decide whether observer's direction shall be read in or calculated.
opt_oosun_read t	1081041	opt_obsdir_read = t - read in angles $\alpha \in [0, 180]$ (measured from the z-axis of the global center-of-mass
		coordinate system, inclination) and $\gamma \in [0,360]$ (measured from the x-axis of the global center-of-mass
		coordinate system, phase angle) from files in_alpha.dat and in_gamma.dat
		opt_obsdir_read = f - Equidistant $\alpha$ , $\gamma$ grid will be calculated based on input options nalpha and
		ngamma.
opt_surface = t	logical	Logical to decide if surface brightness shall be calculated instead of emergent flux profiles.
opt_int2d = f	logical	Logical to decide if surface originatess shall be calculated instead of effective flux profiles.  Logical to decide if the propagation of intensity along a 2D slice trough the computational domain
opt_int2d = 1	logical	shall be calculated instead of emergent flux profiles
ant inal adoubt - f	la si sal	e i
opt_incl_gdark1 = f opt_incl_sdist1 = f	logical logical	Logical to decide if von Zeipel (1924) gravity darkening shall be included for primary object Logical to decide if surface distortion of primary object shall be accounted for
1	0	Logical to decide if surface distortion of primary object shall be accounted for Logical to decide if von Zeipel (1924) gravity darkening shall be included for secondary object
opt_incl_gdark2 = f	logical	
opt_incl_sdist2 = f	logical	Logical to decide if surface distortion of secondary object shall be accounted for
opt_pgrid01 = 'log'	string	Defining the <i>p</i> -grid stratification of the primary object.
opt_rgrid01 = 'log'	string	Defining the <i>r</i> -grid stratification of the primary object.
opt_pgrid02 = 'lin'	string	Defining the <i>p</i> -grid stratification of the secondary object.
opt_rgrid02 = 'lin'	string	Defining the $r$ -grid stratification of the secondary object.
		'lin' – linear stratification
		'log' – logarithmic stratification
		'llog' – log – log stratification
nalpha = 1	integer	Number of $\alpha$ angles to define the directions to the observer
ngamma = 1	integter	Number of $\gamma$ angles to define the directions to the observer
&input_model		Input parameters for the model
vth_fiducial = 1.d2	float	Fiducial thermal velocity $v_{th}^*$ in [km s <sup>-1</sup> ].
&input_surface		Input parameters for surface brightness calculations and calculating intensities along a 2d slice. Will
		be used only if either opt_surface or opt_int2d is set to true
alpha_surface = 1.570796d0	float	The $\alpha$ angle towards the observer.
gamma_surface = 0.d0	float	The $\gamma$ angle towards the observer.
xobs_surface = 0.d0	float	The shift from line center in units of $v_{th}^*$
		Note: In contrast to the single-star version, we here only allow for one surface brightness to be calcu-
		lated at a time.

#### 3.3 cont3Dslab

## 4 Getting started

## 5 Related papers

#### 6 Cite

Depending on the code modules you are using, we would kindly ask you to cite one of the following papers:

- For the ALI scheme using the finite-volume method, please cite Hennicker et al. (2018).
- For the ALI scheme using the short-characteristics method, please cite Hennicker et al. (2020).
- For the formal solution calculating emergent flux profiles or surface brightnesses of single stars, please cite Hennicker et al. (2018) and/or Hennicker et al. (2021).
- For the formal solution calculating emergent flux profiles or surface brightnesses of binary systems, please cite Hennicker et al. (2021).
- For LTE tabulations of occupation numbers, please cite ? (ToDo Luka).

Thank you very much.

For further reading on the ALI method, we refer to Hennicker (2020).

## 7 Developers and contributors

These code modules have been developed in collaboration with: N. Moens, L. Poniatowski., J. Puls, S. Sundqvist. The radiative transfer modules use parts of the GEOMPACK2 library<sup>1</sup> and EISPACK libraries<sup>2</sup> (see Joe (1991)).

## 8 Known problems and solutions

Below, you can find a list of known problems and (possible) solutions:

**OMP is not working:** There are (at least) two possibilities that can cause these problems: OMP\_FLAG is not set in the Makefile (set it to -fopenmp). Alternatively, you might not have set the enivironment variable on your system (in the terminal, simply use export OMP\_NUM\_THREADS=N, with N the number of OMP threads to be used, e.g.,, 12).

Segmentation fault when running in OMP mode: By default, the stacksize for each thread can be very low. Particularly when requiring huge arrays in the formal solution with a lot of grid refinement (e.g., for LDI simulations), the local copies in each thread might run out of stacksize. Probably at the expense of computing efficiency, this problem can be solved by setting the corresponding environment variable: export OMP\_STACKSIZE=10M (or larger if required).

**Comiplation errors of HDF5** Sometimes, a new fortran compiler is not compatible with an old HDF5 version. Then you might want to switch to a later HDF5 release (version 1.10.7 or higher), or downgrade your fortran compiler.

Mac – Illegal Instruction 4: On the Mac, an Illegal Instruction 4 error can occur when static arrays are not properly initialized. To solve this issue, and to still be able to use OpenMP parallelization, please dynamically allocate static arrays, e.g.:

```
real(dp), dimension(nd) :: my_array
becomes
real(dp), dimension(:), allocatable :: my_array
allocate(my_array(nd))
```

## 9 ToDo

• In src/mod\_iline.f90, LTE tables are read in only for 'lte\_tables/Y02800'

https://people.math.sc.edu/Burkardt/f\_src/geompack2/geompack2.html

https://people.sc.fsu.edu/~jburkardt/f77\_src/eispack/eispack.html

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