

A User's Guide to the

Tübingen NLTE Model-Atmosphere Package

TMAP

ATOMS2

SETF2

LTE2

PRO₂

LINE1

LINE1_PROF

and some Auxiliaries

A User's Guide to the

Tübingen NLTE Model-Atmosphere Package TMAP

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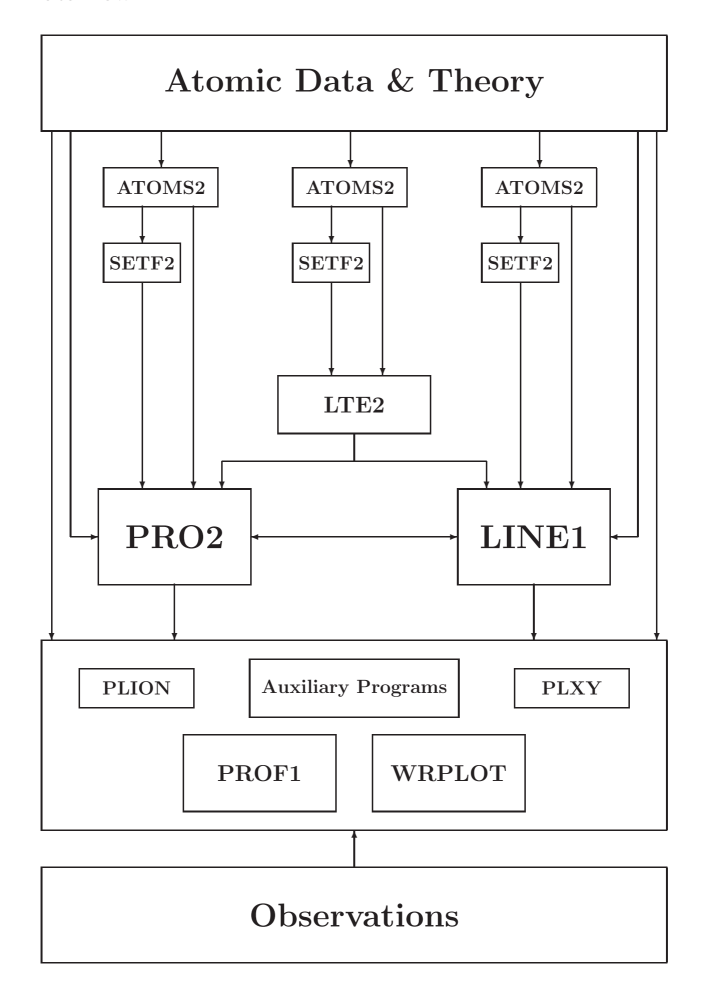
All Users of the Tübingen
NLTE Model-Atmosphere Package
are reminded to be careful
at the creation and / or modification
of the atomic data files.

Although ATOMS2, SETF2, PRO2, and LINE1 check these files partly for consistency, small errors here can have large influence on the resulting model atmospheres.

All programs terminate in the case of fatal errors but can give only warnings when the input is ambiguous. The neglection of these warnings may be fatal!

For *PRO2* and *LINE1* output options are existing to print the model atoms and atomic data read by these programs. They should be used for checks — not only at the first time when the program is running with new or modified data!

Data flow



Overview 3

0 Overview

In the following the Tübingen NLTE Model-Atmosphere Package (TMAP) is described, which is based on the approximated lambda iteration (ALI) method. It consists mainly out of a "LTE" model atmosphere program (LTE2) for the calculation of start models, a NLTE program (PRO2), a NLTE line formation program (LINE1) and a plot program (PROF1). To start the programs you need the following:

- 1. an executable *program* of the most recent program version. Details can be found in Sect. 1.
- 2. a model atom file with all model atoms and atomic data. This file is created by the user and partly checked and automatically filled up by the program ATOMS2 (Sect. 2).
- 3. a frequency grid file, which contains the frequency points, the respective quadrature weights, and informations about every frequency point. It is created by the program SETF2. Details can be found in Section 3.
- 4. a *start model*. It can be either a "LTE" start model or a model which has been created by *PRO2* or *LINE1*. Further informations are summarized in Section 4.
- 5. commands for the program control (Sect. 5.4) and print commands (Sect. 5.5).

1 Creation of the Executables

All programs of the Tübingen NLTE Model-Atmosphere Package (*TMAP*)were developed on CRAY computers. They can generally be installed on all other computers. For the substitution of CRAY specific parts or system library routines, there exists a collection of own subroutines.

For the use of a new atomic data or frequency grid files it might be necessary to change the PARAMETER statements for the DIMENSION of various arrays. Subsequently the program has to be compiled and loaded. Shell scripts for compilation are available. *Please contact rauch@astro.unituebingen.de for further instructions.*

All PARAMETER statements are collected in input files which are named PARAND, PARA, PARA1, and PARA3. All PARAMETER signed with "*" are used by PRO2 only, all with "*" by LINE1 only, the rest is valid for both programs.

PARANA

```
NA quadrature points for angular integration (default 3). PRO2 allows NA \in [3, 5]. LINE1_PROF allows NA \in [3, 4, 5, 6, 7, 8, 9, 10, 12, 16, 20, 24, 32].
```

PARAND

```
ND depth points (default 90)
```

An investigation on the necessary number of depth points has shown that the computational time decreases (of course) for ND < 90 but the resulting line profiles are more narrow and the line cores are less deep. The line profiles converge at ND $\stackrel{>}{\scriptstyle \sim}$ 70. An 1% error is reached at ND $\stackrel{>}{\scriptstyle \sim}$ 90. For much higher ND, the computational time increases tremendously and the models a less numerical stable.

PARA

LTEMAX LTE levels

NATOMAX elements

NFMAX frequency points

NIONMAX ionization stages

NLATOMS non-LTE levels (of one element)

NLMAX non-LTE levels (total of all elements)

PARA1

NRBBMAX transitions radiative bound-bound

NRBFMAX transitions radiative bound-free

NRFFMAX transitions radiative free-free

NRLLMAX transitions **rbb** for "sample" cross-sections within one level band (complex ions) (Sect. 2.2)

NRLUMAX transitions **rbb** for "sample" cross-sections (Sect. 2.2)

NCBBMAX transitions collisional bound-bound, (only electron collisions)

NCBFMAX transitions collisional bound-free, (only electron collisions)

NCBXMAX transitions collisional bound-extra, (only electron collisions)

as well as the same PARAMETER with a D instead of X at the end (NXXXMAD): maximum number of input values in the atomic data file for the transition XXX (e.g. NCBBMAD)

Exception: NRBBMAD, either like above or – if larger – maximum number of frequency points in a line transition

NRBFMAW* DIMENSION of the array SIGBF

NRBLMAX* blends of a line for which a detailed line profile is calculated

NRLFMAX* lines to select in a single job for the calculation of their profile

PARA3

NRDIMAX transitions radiative dielectronic bound-free

NRDIMAD (analogous PARA1)

 $NSIG^*$ DIMENSION of the array SIGBF (value NRBFMAW (= NSIG) is given by SETF2 and $LINE1_PROF$)

In the case that the PARAMETER limit is exceeded, e.g. the number of NLTE-levels in the atomic-data file is higher than NLMAX, the programs stop with an error message, partly only in the log file (STOP '....'). Working with model atoms much below the PARAMETER limits is a waste of core memory and I/O time. Take care that the PARAMETER values do not create bank conflicts, i.e. avoid 2^n values!

2 Creation of Atomic Data Files

All programs expect an atomic data file, which contains model atoms and atomic data. It is firstly created by the user (Sect. 2.7) and then processed by the program *ATOMS2*.

The model ions of one element have to be inserted in increasing order while any order of elements is accepted.

The processing of the atomic data file is directed by *keywords* which meanings are explained in the following. All **keywords** as well as all **strings** have to be inserted flush left (e.g. level names, Sect. 2.1). All **numerical data** can be written format free. *Attention: all input lines have a maximum length of 80 characters!*

2.1 Level Names (TMAP Code)

The levels are named using an A10 string. The element code is at the beginning (e.g. HE) and the following (one or two) numbers indicate the ionization stage (e.g. HE2). This is mandatory. The next numbers indicate the principal quantum number of the level (e.g. HE26). For levels with a complicate configuration, the electron system and the magnetic quantum number may be added (e.g. O42P 2PO). Singly excited levels are marked by a "'," doubly excited by a "'," directly before the electron system indication (e.g. N33S' 2PO). In the case of explicitly considered multiplet splitting the angular momentum is added at the end (e.g. C452D5/2).

2.2 List of Keywords for the Atomic Data File

O ATOM L LTE RBB RBF RDI RLL RLU CBB CBF CBX TL

2.3 Description of the Keywords

The expression "card" which is frequently used in the following is a relic from the PUNCH era and is used for historical reasons.

```
keyword.
```

DB

```
comment, not written to atomic data file
```

```
keyword 0 (Null)
```

necessary to end the validity of some keywords (e.g. L, LTE, ...)

keyword ATOM

introduces a new element. The following card indicates:

- 1. the chemical abbreviation (FORMAT A2),
- 2. charge of the lowest ionization stage (in e^-) in model atom,
- 3. atomic weight (in AMU).

Example:

MOTA

HE 0 4.0026

means: all following cards describe an helium model atom, starting with neutral helium He I, the atomic weight is 4.0026 AMU.

keyword L

introduces non-LTE levels. All cards following this keyword indicate:

- 1. level name (FORMAT A10, Sect. 2.1),
- 2. name of ground state of the following ionization stage (FORMAT A10, Sect. 2.1), if **no** parent exists: NONE.
- 3. energy to the ionization limit in Hz,
- 4. statistical weight, if **no** parent exists: 1.0.

Example:

L

HE26_{LULULULU}HE31_{LULULULU}3.6548882425E+14 72

levels of an ionization stage are expected **in increasing energetic order** (from the ground state). The keyword 0 completes the list of NLTE levels for this ionization stage.

keyword LTE

introduces LTE levels analogously to the keyword **L**. Do not forget to complete the list with the keyword **0**. Important: some formulae for the free-free opacity (Sect. A) expect at least one LTE level in the respective ion. Attention: this is **not checked** by the program.

Radiative and collisional transitions are introduced by:

RBB radiative bb

RBF radiative bf

RDI radiative dielectronic bf

RLL radiative bb for "sample" cross-section within one level band (complex ions)

RLU radiative bb for "sample" cross-section

CBB collisional bb

CBF collisional bf

CBX collisional from NLTE to LTE levels

All lists following these keywords have to be completed by the keyword 0. A card which follows one of the keywords (except RDI) indicates

- 1. lower level (A10),
- 2. upper level (A10),
- 3. formula number for the calculation of the cross-section (Sect. A),
- 4. number n of following input data,
- 5. n input data for the calculation of the cross-section.

Example:

```
RBB
```

```
H11<sub>____</sub>H13<sub>___</sub>1 1 0.0791
0
```

This is the line transition Ly_{β} , cross-section calculation with formula No. 1, one input number (0.0791).

For RDI transitions a third level is introduced between lower and upper level which is the upper level of the stabilizing transition and **has to be** introduced before as a LTE level.

For RLU and RLL transitions the level names are followed by the name of the file which contains the "sample" cross-section.

Example:

```
RLU
```

keyword RFF

introduces radiative free-free transitions. The following card indicates:

- 1. the ionization stage (A10, chemical element abbreviation + ionization stage, e.g. HE2),
- 2. formula number for the calculation of the cross-section (Attention: Def. LTE!)
- 3. number n of the following input data,
- 4. *n* input data.

keyword TL

The card which follows this keyword indicates the "line temperature" T_{line} for all following RBB transitions. This is necessary to change the temperature used for the calculation of the Doppler width (default $T_{\text{line}} = \frac{3}{4}T_{\text{eff}}$) in the creation of the frequency grid (Sect. 3). The default is again valid after

TL

0

keyword DB

introduces an explicit frequency grid for all following RBB transitions. The following card indicate:

- 1. number n of frequency points of the line,
- 2. n frequency points (distance from line center in Doppler widths!).

Example:

DB 7 -3 -2 -1 0 1 2 3

If the first frequency point is 0.0, only "half" lines are created.

This grid is valid until the next keyword DB appears. The default is valid after

DB 0

Note: In the Tübingen Model-Atom Database (Sect. 2.4), standard DB values are given that are about the maximum needed at low effective temperatures and high surface gravities. These values have to be altered at least for lower surface gravities. A test calculation with PRO2 will show the line widths. Use, e.g., $HI\lambda 1215 \text{ Å}$, and reduce the maximum DB values. In case that they are too low, "steps" in the flux level are prominent in the line wings. (The Doppler width of a specific line can be calculated by /home/rauch/bimod/doppler.Linux_x64.)

The DB adjustment results in a lower number of frequency points and, thus, faster model calculation.

A good compromise may also be to replace

```
DB
33
-2048.0 -1024.0
                    -512.0
                              -256.0
 -128.0
            -64.0
                     -32.0
                               -16.0
   -8.0
             -4.0
                       -2.0
                                -1.0
   -0.5
             -0.3
                       -0.2
                                -0.1
0.0
    0.1
              0.2
                        0.3
                                 0.5
    1.0
              2.0
                        4.0
                                 8.0
   16.0
             32.0
                       64.0
                               128.0
  256.0
            512.0
                    1024.0
                              2048.0
```

by the insertion of a grid of additional frequency points using, e.g.,

```
/home/rauch/bimod/prep\_conts\_man.Linux\_x64 > F\_BASE << eos 1175 1255 0.5 eos
```

and a reduced DB that represents the line core much finer than the base grid (for F_BASE see Sect. 3)

```
DB
15
-2048.0
                        -2.0
                                 -1.0
             -0.3
                        -0.2
                                 -0.1
    -0.5
0.0
                        0.3
     0.1
               0.2
                                  0.5
     1.0
               2.0
                                   2048.0
```

2.4 TMAD

The Tübingen model-atom database TMAD (http://astro.uni-tuebingen.de/~TMAD) provides ready-to-use model atoms in TMAP format.

2.4.1 Model calculations (PRO2)

TMAD provides, in general, model atoms at maximum size, i.e. using all data from standard atomic-data resources. For model calculations, model ions can be reduced typically to 15 to 20 NLTE levels (Sect. 2.5, cf. Jahn et al. 2007) to avoid unreasonable computation times. In addition, the input for the DB keyword (Sect. 2.5) has to be adjusted to avoid calculations of line profile very far off the line centers where the line absorption is negligible. TMAD uses a standard of 2400 Doppler widths which may be necessary only for models with very high surface gravities.

2.4.2 Line-formation calculations (PRO2)

Models from Sect. 2.4.1 may not include all levels, e.g. for C IV lines in the optical wavelength range around 4660 Å. Thus, a subsequent line-formation calculation has to be performed with fixed atmospheric structure, i.e. only NLTE occupation numbers for the atomic levels are calculated. It may not be useful to extend all ions and to consider all levels given in TMAD. The cookbook says, "for all ions for which lines are identified, include five more levels than the highest from which strategic lines are arising".

In case of the C IV $\lambda\lambda4660\,\text{Å}$ / He II $\lambda4685\,\text{Å}$ absorption trough in PG 1159 stars, it is sufficient to extend C IV only for a line-formation calculation, even in models that consider all species from H to Ni.

LINE1 and LINE1_PROF provide the possibility to collect occupations numbers of newly considered levels from different models Sect. 8.3, e.g. if five line-formation calculations were performed (based on the same model) with individually extended CIV, OV, OVI, NeVI, and ArVI. This saves an enormous amount of CPU time.

2.4.3 Spectra calculations (LINE1_PROF)

To calculate the emergent synthetic spectrum, a formal solution is performed with LINE1_PROF. TMAD provides model atoms that account for fine-structure splitting of the atomic levels. These have to be the same like the unsplitted levels used for the line-formation calculations. (Sect.2.4.2).

2.5 ATOMS2

The interactively created atomic data file is — if designed following the instructions in Sect. 2 — ready, i.e. it can be used for the creation of a frequency grid as well as for model atmosphere calculations. However, it is highly recommended to process it with the program ATOMS2, which is able to detect a lot of errors and gives warnings. In case of fatal errors it even terminates. There exist options which make ATOMS2 create automatically model ions or fill up them.

The program *ATOMS2* is available at Tübingen's PC/Workstation cluster in its latest version: /home/rauch/bimod/atoms2.Linux_x64.

An atomic data file (ATOMS) created by ATOMS2 is unambiguously defined by:

- the input file (here: *OPTIONEN*)
- the interactively created atomic data file (here: ATOMIN)

In the input file OPTIONEN one can write directives for ATOMS2. ATOMS2 expects OPTIONEN and ATOMIN subsequently as input, OPTIONEN has to be finished in any case with END_OPTIONS,

even if none of the following options is given:

$$AUTO_{\square}ION_{\square}H_{\square\square\square\square}I$$
 in il ir

This option creates a complete H model atom. in is the principal quantum number of the highest H I NLTE level, il is the principal quantum number of the highest H I LTE level, ir is the principal quantum number of the highest H I NLTE levels up to which all RBB transitions are considered. (The conditions il > in and $ir \le in$ are fulfilled.) In case that ir is negative, only transitions from ATOMIN up to the NLTE level ir are used. (This is necessary if ATOMIN contains information about line broadening, default is Doppler line broadening). In the case that ir is positive, ATOMIN does not need to contain any H model atom.

same like $\mathtt{AUTO}_{\square}\mathtt{ION}_{\square}\mathtt{H}_{\square\square\square\square}\mathtt{I}$ (see above) but mandatory; in the case that ATOMIN contains a He is model ion, ATOMS2 creates He ii and He iii model ions, otherwise a complete He model atom.

CBB-AUTO-FILL

If this option is given, missing CBB transitions are automatically inserted and possible errors in ATOMIN corrected — as far as ATOMS2 is able to detect them. The output of ATOMS2 has to be checked for respective error messages or warnings.

same like CBB-AUTO-FILL; possible errors are only reported in the output but not corrected.

If this option is given, all CBB transitions are disregarded. This ...-AUTO-FILL option is only useful for the detailed calculation line profiles. This option is automatically set, if the option LINEFORMATION ... is given (see below).

Analogously to the CBB-AUTO-FILL ... options, the following options are valid:

```
CBF-AUTO-FILL
```

(for unknown cross-sections, hydrogen-like values are automatically inserted)

(for OpacityProject cross-sections, hydrogen-like values are automatically inserted)

CBX-AUTO-FILL

 $CBX-AUTO-FILL_{||||}(IGNORE)$

CBX-AUTO-FILL (NONE)

RBB-AUTO-FILL (NONE)

RBB-AUTO-FILL (H1 H1)

 $\texttt{RBB-AUTO-FILL}_{\sqcup\sqcup}(\c_{|\;|\;|}\texttt{HE2}_{|\;|})$

```
RBF-AUTO-FILL ( NOOP) (for OpacityProject cross-sections, hydrogen-like values are automatically inserted) (set CBF-AUTO-FILL ( NOOP) also) (RDI-AUTO-FILL ( NONE) ( NONE) ( NONE) ( NONE) ( NONE) ( NONE) ( NONE)
```

```
LINEFORMATION, -, RBB-INTERVALL=[,,,1000,,,,7000],,,10N=H1
```

This option can be set for line-profile calculations in order to restrict number of line transitions in the atomic data file ATOMS to a selected interval (here: [1000, 7000] Å). If this shall be valid for all (line profiles), ION=NON has to be inserted. To insert all line transitions — i.e. even outside the given interval — of a selected ion in ATOMS (line formation), the TMAP code (Sect. 2.1, here: H1) has to be given.

```
If this option is given, all C.. -AUTO-ION _{\square\square} (_{\square\square}NONE) options and the RDI _{\square}-AUTO-ION _{\square\square} (_{\square\square}NONE) option are automatically set.
```

```
LIST OF ALL RBB, ION:
```

This option creates a table of all line transitions of the selected ion under consideration of a possible LINEFORMATION ... option with its interval restriction.

END, OPTIONS

This option finishes the file *OPTIONEN* and is necessary if one of the other options has been given.

The program ATOMS2 creates a large informative output which includes generally the atomic data file as created by ATOMS2, informations about wavelengths of levels, thresholds, line transitions, and some statistics. Excited levels are marked in the output with "**", all transitions which were inserted following the ...AUTO-FILL ... options are marked with "AF".

All parameters which are valid for the created atomic data file ATOMS (Sect. 1), are summarized in a table at the end of the output. Under unix a

```
grep para joutput filename;
```

extracts them from the output file.

2.6 TIRO

The Tübingen Iron-Group Opacity (TIRO) service creates atomic data files and cross-section data for radiative bound-bound and bound-free transitions of iron-group elements (calcium, scandium, titanium, vanadium, chromium, manganese, iron, cobalt, and nickel). It is based on the program IrOnIc that was developed at Tübingen. TIRO enables the VO user to consider iron-group elements in model-atmosphere calculations easily, in various ways, and without spending own

calculation time for the creation of the necessary input data. It is controlled via web interface (http://astro.uni-tuebingen.de/~TIRO) in which the following inputs have to be given.

- The name, institute, and email address of the user have to be given to inform the user about the status of the process and the location of the results.
- The resulting line profiles depend on the line temperature $T_{\rm L}$. $T_{\rm L}$ is $3/4~T_{\rm eff}$ and corresponds to the mean temperature in the lineforming region.
- A frequency grid can be uploaded. The corresponding file has to be written in plain text with monotonically increasing frequency points and must be in *TMAP* (http://astro.uni-tuebingen.de/~TMAP) format. Alternatively, a start and end wavelength as well as a spacing or maximum number of points can be chosen. The frequency grid is then created automatically by *TIRO*. The calculations are performed on this frequency grid or already calculated cross-sections from the database are interpolated to this frequency grid.
- It is possible to retrieve data for all ionization stages up to IX. The requested ionization stages can be chosen.
- Cross-sections for the iron-group elements can be created considering all or some of them individually or generically. When the individual option is chosen, an easy adjustment of the abundances in the model atmosphere calculation is possible. A generic model atom consists of selected iron-group elements. In this case, the given abundance ratios are fixed in the resulting model atom. For both cases the line type of the underlaying atomic data can be selected. They are taken from Kurucz' line lists and can be chosen between lines with measured wavelengths (POS) or measured and calculated wavelengths in addition (LIN).

After submitting the data, the given parameters are stored in a request file. TIRO checks regularly if requests are waiting and processes them one after the other. The user is informed via email when the handling of the data starts. The resulting files are stored in a compressed tar archive that is accessible via a wget command. The user is informed via email about its location. The files for bound-bound transitions contain a table with frequencies in the first column, cross-sections in the second (calculated for electron density 0) and in the third column (calculated for electron density $10^{16}/\text{cm}^3$). The corresponding files for bound-free transitions include a table with frequencies in the first and cross-sections in the second column.

2.7 Auxiliary Programs

Some programs are currently available which help to design model atoms etc. All these programs work interactive and are (more or less:-(...) self-explaining.

2.7.1 FORMEL4

The programs PRO2 and LINE1 can consider – besides pure Doppler line broadening – the quadratic ("formula 3") and the linear ("formula 4" — this gives the name of the program ...) Stark effect for the line broadening. Both formulas need some input data which has to be includes in the atomic data file ATOMS. (Sect. 2).

The program FORMEL4 (/home/rauch/bin/formel4) calculates these data (mainly the classical damping constant and the effective quantum number) from an existing atomic data file.

The calculated data is saved into the files FORMEL4.DAT and FORMEL4.RBB. FORMEL4.RBB can be used the replace the respective section of the atomic data file which was used to calculate the data.

FORMEL4 inserts also the vacuum wavelengths, starting with the keyword WAVELENGTH: in columns 81-91. The wavelength values may be the replaced by measured wavelengths in order to shift those lines which arise from levels with uncertain energies to the correct wavelengths. A subsequent run (necessary!) of SETF2 on the modified ATOMS yield a frequency grid with these lines at the correct wavelengths.

2.7.2 LEVEL

The level energies in literature are commonly given in cm^{-1} , measured from the ground state of their ionization state. In the Tübingen NLTE Model-Atmosphere Package (TMAP), the level energies given are the difference to the next ionization limit, i.e. the energy which is necessary to ionize into the ground state of the next ionization stage. The program LEVEL simply transforms these energies.

The program LEVEL (/home/rauch/bin/level) runs interactively. It is possible to calculate

- energies of single levels
- energies of combined levels

Furthermore it is possible to combine

• levels from an existing atomic data file

The data is saved into LEVEL.DAT and (for the combination of levels) into XX.ENTARTET (XX is the principal quantum number of the combined levels).

2.7.3 MKIDENT_FROM_KURUCZ

Due to our statistical approach to create atomic data files from Kurucz's data (Kurucz 1991, 2009, 2011) of elements with atomic numbers $Z \geq 20$, the wavelengths of individual lines can only be read from Kurucz's data files. The two procedures

/home/rauch/tools/lineidentification_LIN.bat and

/home/rauch/tools/lineidentification_POS.bat

for Kurucz's data files with theoretical and laboratory measured lines (.LIN) and with positively identified lines (.POS), respectively) may be used to create

\IDENT <wavlength> <ion>

cards that can be used by WRPLOT (Sect. 11). An explanation how to use these procedures is found in their headers.

2.7.4 MULTIPLET

During the calculation of the model atmospheres, multiplets are considered with a combined level. For the subsequent detailed line profile calculation, it is necessary to split these multiplets into single components. While (in most cases) the level energies can be taken from literature, there are only rudimentary informations about the oscillator strengths of the single components. The program MULTIPLET (/home/rauch/bin/multiplet) splits up oscillator strengths of doublets, triplets, and quartets under the assumption of LS coupling.

The calculated data is saved into MULTIPLET.DAT.

2.7.5 **SEATON**

For some RBF cross-sections of levels of the ions He I, C I – C IV, N I – N V, O I – O III, Ne I – Ne II, Mg I – Mg II, Si I – Si IV etc., tables have been calculated by Hofs" a"s. These can be evaluated with

the program SEATON (). One gets input parameter for the Seaton formula (Sect. A.4). For those levels which are not included in the tables, SEATON can calculate a hydrogen-like cross-section.

2.7.6 SGF

To reduce the pixel-to-pixel variation (or the noise) of observations, a low-pass filter may be used. The program SGF (/home/rauch/bin/sgf) processes data following Savitzky & Golay (1964). Two parameters, np and m have to be given. np is the number of a subset of the data which is used for a fit. m is the degree of the polynomial used in the least-squares fit method. m has to be carefully chosen - a too-high value smoothes out real spectral features.

3 Creation of Frequency Grids

The program SETF2 creates the binary file FGRID which contains the frequency grid used by LTE2, PRO2, and LINE1.

SETF2 processes data from the atomic data file ATOMS. Attention: SETF2 has not the capabilities to detect errors like ATOMS2! For the creation of a frequency grid SETF2 uses only data which belongs to the keywords L, RBF und RBB (Sect. 2.2) and checks these for errors.

Thus, it is indispensable to transform every new or modified, interactively created atomic data file with the program ATOMS2 in order to check for errors and inner consistency.

The program SETF2 is available for all users at Tübingen's PC cluster in its latest version: $/\text{home/rauch/bimod/setf2.Linux}_{<}X86|P4|x64>$.

A frequency grid is unambiguously defined by its input files ATOMS, DATEN, $CONTS_MAN$, F_BASE , and POS_LIST . All frequency points are clearly named (TYP) in the frequency grid (see below.).

ATOMS

is the atomic data file (Sect. 2).

DATEN

This file contains:

- the effective temperature T_{eff} ,
- the output parameter,
- the weight parameter $\nu_{\rm W}$,
- the line parameter,
- the EUV parameters EUVmax und EUVnumber,
- f-value limit.

The effective temperature $T_{\rm eff}$ is necessary for the calculation of the frequency discretization within a line transition (central frequency ν_0). The position of the frequency points is depending on the Doppler width $\Delta\nu_{\rm Doppler} = \frac{\nu_0}{c} \sqrt{\frac{2kT}{m_{\rm ATOM}}}$. This default value can individually be substituted by the keywords TL in ATOMS.

If the "RBB" record in *ATOMS* contains the keyword WAVELENGTH: in columns 81-91, *SETF2* uses the value directly following this keyword for the central wavelength of the respective line transition. The program *FORMEL4* can preset this value which may be replaced or inserted (with the keyword!) manually in order to shift the line to the specified wavelength.

Two edge frequency points are inserted for every bound-free transition in ATOMS (keyword RBF): this is necessary because the ionization into excited levels is considered in detail by PRO2 und LINE1. These frequency points are set at the ionization energy ν_0 of the respective level and at $\nu_0 + 10^{-12} \cdot \nu_0$. These points are used to split up the complete frequency grid in single integration intervals.

For all line transitions in *ATOMS* a particular number of frequency points is inserted. Their order is controlled by the *line parameter*. If nothing else is defined in *ATOMS* (keyword DB), the following arrangements are automatically set:

In the case that the frequency points which are inserted for a line transition overlap with those of another line or with edge frequency points, the overlapping lines have to have a symmetric arrangement of their frequency points ("full lines" — like inserted if the *line parameter* is set to 1, Tab.1)

The choice of line parameter 0 is generally possible for all other line transitions under the

line parameter frequency points $(\nu - \nu_0)/\Delta\nu_{\text{Doppler}}$ comment 0 5 $0, \frac{2}{3}, \frac{4}{3}, 2, \frac{8}{3}$ universal 1 9 $-\frac{8}{3}, -2, -\frac{4}{3}, -\frac{2}{3}, 0, \frac{2}{3}, \frac{4}{3}, 2, \frac{8}{3}$ universal -1 5 like 0, in case of line overlaps like 1 universal	200010 1. 2110	verce Paracritector engre	r delegate ir equelley perme erromoune	0 101 11110 010011010
1 9 $-\frac{8}{3}, -2, -\frac{4}{3}, -\frac{2}{3}, 0, \frac{2}{3}, \frac{4}{3}, 2, \frac{8}{3}$ universal	line parameter	frequency points	$(\nu - \nu_0)/\Delta \nu_{\mathrm{Doppler}}$	comment
-3 $0, \frac{4}{3}, \frac{8}{3}$ pre-iteration		5 9 5 3	$-\frac{8}{3}$, -2 , $-\frac{4}{3}$, $-\frac{2}{3}$, 0 , $\frac{2}{3}$, $\frac{4}{3}$, 2 , $\frac{8}{3}$ like 0, in case of line overlaps like 1	universal universal

Table 1: The *line parameter* and default frequency point arrangements for line transitions

premise that the flux gradient is almost constant over the full width of the line. This kind of frequency discretization within a line transition is named "half line".

An automatical differentiation is made if the *line parameter* is set to -1. This has the great advantage that fewer frequency points are inserted in the grid, compared to *line parameter* 1.

The *line parameter* -3 is equivalent to *line parameter* -1 but inserts only three points (five in case of overlap) for a line. This saves memory and computational time but is somewhat unrealistic because the line profile is not well represented by this frequency point arrangement. At least a subsequent iteration with a more detailed frequency grid (e.g. created by *SETF2* from the same atomic data file *ATOMS* but with the *line parameter* -1) is necessary.

The f-value limit can be chosen to represent lines with f-values smaller equal than the f-value limit analogously to $line\ parameter$ -3, independent from DB values in ATOMS. This reduces slightly the number of frequency points in FGRID.

The frequency grid contains by default the interval $1.0 \cdot 10^{12} - 2.99792458 \cdot 10^{17}$ Hz (= 1 Å, Tab.3). The optional parameter EUVmax allows to change the maximum: in the case that EUVmax is larger than $2.99792458 \cdot 10^{17}$ Hz, EUVnumber indicates the number of frequency points which are inserted by SETF2 in the range $1.0 \cdot 10^{17} - EUVmax$ Hz; in the case that EUVmax is smaller that $2.99792458 \cdot 10^{17}$ Hz, EUVnumber is disregarded.

CONTS_MAN

This optional file contains additional frequency points (given in Å or Hz)

In this input file frequency points can be requested to insert which are important for the evaluation of the models, e.g. in the case of the emergent flux and the calculation of specific colors. Doubly requested points are disregarded . . .

CONTS_MAN can be created using

/home/rauch/bimod/prep_contsman.Linux_<X86 |P4|x64> > CONTS_MAN << eos min max < step / resolving power> eos

for wavelength or frequency points. In case that the third argument is ≥ 100 , prep_contsman assumes it to be the resolving power. prep_contsman accepts more than one input line.

F_BASE

This optional file contains additional frequency points (given in Å or Hz)

If the file F_BASE exists and contains valid input data, all requested points are created as base grid in the interval $\nu_{\min}^{\text{base}} - \nu_{\max}^{\text{base}}$. The largest base grid can actually contain 200 000 frequency points. Subsequently, SETF2 creates (like described above) the "normal" frequency grid based on ATOMS and DATEN. The line parameter is automatically set to 1.

In contrast to frequency points requested in *CONTS_MAN* which are definitely present in the created *FGRID*, all base grid points are eliminated if "normal" frequency points (including points

requested in $CONTS_MAN$) are found in the interval $\left[\frac{1}{2}\left(\nu_{i-1}^{base} + \nu_{i}^{base}\right), \frac{1}{2}\left(\nu_{i}^{base} + \nu_{i+1}^{base}\right)\right]$ Hz. F_BASE can be created using prep_contsman, see above.

POS_LIST

This optional file contains additional frequency points (given in Å or Hz)

It should be created by /home/rauch/tools/line_identification_POS.job (edited for the required elements and ionic species) for Kurucz POS lines only.

SETF2 reads the line centers from **POS_LIST** and creates for each line nine frequency points $(\Delta \lambda/\mathring{A} = -0.005, -0.003, -0.002, -0.001, 0.000, 0.001, 0.002, 0.003, 0.005)$. This is necessary to avoid artificial "fading" of iron-group lines by a later convolution of the synthetic spectrum with a rotational of instrumental profile.

After the creation of the frequency grid in accordance with ATOMS, DATEN, CONTS_MAN, F_BASE, POS_LIST the IR, UV, and EUV range and the intervals between the lines and edge are automatically completed.

Every frequency point is unambiguously named in the frequency grid with a CHARACTER*50 string: In detail the fields have the following meaning:

field 1:

- CONT for all continuum points, which were set in accordance to ATOMS, DATEN, $CONTS_MAN$, and F_BASE
- C/ED for all continuum points, which were automatically created, e.g. in order to separate integration intervals
- EDGE for all continuum points, which were inserted to represent the bound-free transitions in *ATOMS* (see above)
- LINE for all line transition points

field 2:

- (HALF) for all line transition points, "half lines" (see above)
- (FULL) for all line transition points, "full lines" (see above)
- $_{\square}$ RED $_{\square\square}$ for all continuum points at the end of integration intervals (in general from one bound-free transition to another)
- $_{\square}$ BLUE $_{\square}$ $_{\square}$ RED $_{\square\square}$ (RED and BLUE indicate the location of the point relatively to the threshold/edge: BLUE is set exactly at the threshold frequency, RED is set at longer wavelength, i.e. "redwards")

field 3:

• Lilling ** for threshold frequency points for levels ionizing into excited levels

field 4:

- AUTO_ADD___ for all automatically inserted continuum frequency points (to fill-up too-large intervals)
- AUTO_IR_ $\square\square\square$ for all automatically inserted continuum frequency points (to fill-up to the IR limit of $1.0 \cdot 10^{12} \text{Hz}$)
- AUTO_UV for all automatically inserted continuum frequency points (to fill-up to the EUV limit of 2.99792458 · 10¹⁷Hz)
- BASE ____ for all inserted (and not eliminated see above) continuum frequency points optionally requested in F_BASE
- \bullet MANUELL for all inserted continuum frequency points optionally requested in $CONTS_MAN$
- HE26 for all edge and line transition frequency points lower level in the *TMAP* code (Sect. 2.1, here: He II, level with principal quantum number 6.

$\underline{\text{field 5}}$:

• HE27 for all edge and line transition frequency points upper level in the *TMAP* code (Sect. 2.1, here: He II, level with principal quantum number 7

field 6:

For all line transition points the position of the point in the line is indicated:

- CONT/RED
- CENTER
- CONT/BLUE

For every frequency point a quadrature weight is calculated corresponding to the weight and the *line parameter*. The following points are considered:

- integration intervals do not include a complete edge, i.e. EDGE/RED and EDGE/BLUE of the same threshold
- the quadrature weights are calculated following the trapezoidal law for $\nu < \nu_{\rm W}$ (weight parameter, see above) and following the Simpson law for $\nu_{\rm W} < \nu$. The Simpson law is necessary because the flux is exponentially decreasing for $\nu > \nu_{\rm W}$ and the trapezoid law is not sufficient. In principle, all weights can be calculated following the Simpson law SETF2 will then insert a few more frequency points.
- "half lines" have corrected weights
- In the case of line overlaps "full lines" have to be considered (automatically: *line parameter* 1 or better -1, or keyword DB in *ATOMS*)

The output of SETF2 should be controlled in detail. SETF2 comments all operations and gives warnings in case of ambiguity. In the case of fatal errors, SETF2 terminates.

The quadrature weights require special attention: the check sums of all single intervals as well as of the complete frequency grid have to be exactly 1. If this is not the case, it is possible

that negative quadrature weights appear ("half lines" in cases were "full lines" are necessary). In order to correct for this, the atomic data file ATOMS has to be modified (keyword DB) or the line parameter has to be set to -1 (this automatically corrects these cases).

Attention: A frequency grid with wrong quadrature weights will cause an abort in programs like PRO2 or LINE1.

The size of the output is controlled by the *output parameter*:

- PRINT CHECK reduces the output to a minimum, only warnings, fatal errors, and information about the quadrature weights are displayed.
- ullet PRINT gives a complete output, incl. the complete frequency grid. In both cases the parameter NFMAX, NRBBMAX, NRBBMAD, and NRBFMAW (Sect. 1) are printed, which are valid for ATOMS and FGRID. Under unix a

grep para joutput filename;

extracts them from the output file.

• PRINT ASCII FGRID is created formatted.

In case that the input atomic-data file was created by ATOMS2 for line-profile calculations (Sect. 2), SETF2 creates two files, ION_USER and $LINIEN_USER$, that may be used as input files (ION and LINIEN) for $LINE1_PROF$ (Sect. 9). $LINIEN_USER$ contains a line closest to the center of the RBB interval (ATOMS2, Sect. 2) and a respective BLENDRANGE to cover the complete RBB interval. ION_USER contains the proper ion according to $LINIEN_USER$.

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4 Start Models

Start models are necessary for the programs PRO2 and LINE1. They can come from different sources. Within the Tübingen NLTE Model-Atmosphere Package (TMAP)all models can be used as start models. It is distinguished between two types of start models.

4.1 PRO2 and LINE1 Models

A model which is calculated by PRO2 or LINE1 can be used as start model. The convergent models (with the same parameters, same ATOMS and FGRID) from both programs should be identical.

New NLTE-levels, i.e. levels that are not found in the start model, are set to LTE occupation numbers. This is valid especially for completely new elements and can be a problem in the case of higher elemental abundances due to the inconsistency in the conservation of particles . . .

For the detailed calculation of line profiles, automatic multiplet splitting is optional in *LINE1* for all C IV, N V, O VI levels and a wide variety of C III, N IV, and O V levels. The list of multiplets is permanently updated.

4.2 "LTE" Models

The program LTE2 produces "LTE" models (without any consideration of line blanketing) within a large range of photospheric parameters. The created model has PRO2 format and is a start model for PRO2 or LINE1.

4.3 Calculation of a LTE2 Model

The program LTE2 calculates models in gray approximation. It uses the same atomic data (ATOMS, Sect. 2) and frequency grid files (FGRID, Sect. 3) like PRO2 and LINE1. No line blanketing is considered. LTE2 is controlled via the input file DATEN with a number of options (Sect. 4.3.2).

4.3.1 The Method

The program LTE2 calculates a model atmosphere based on the requested effective temperature T_{eff} , surface gravity g, chemical composition, and the files ATOMS and FGRID.

An equidistant $\log \tau$ scale (τ is the Rosseland optical depth) with ND depth points (Sect. 1) is created on which start values for the temperature stratification are calculated. This $\log \tau$ scale can be manipulated by some options (see below).

With the basic assumption of LTE,

$$S_{
u} \equiv B_{
u}(T),$$

the mean intensity of the radiation field is

$$J(au) = S(au) = B\left[T(au)
ight] = rac{\pi}{\sigma_B} \cdot T^4$$

i.e. the flux is constant $(\frac{dH}{d\tau} = J - S \equiv 0)$. Thus, also the 1st moment of the radiation field $\frac{dK}{d\tau} = H$ is constant and yields the exact integral

$$K(au) = H \cdot au + c = rac{1}{4} F \cdot au + c$$

From this, we get $K(\tau) \to \frac{1}{4}F \cdot \tau$ for $\tau \gg 1$. $K(\tau) = \frac{1}{3}J(\tau)$ for $\tau \gg 1$ is the general solution for the intensity

$$J(au) = rac{3}{4} F\left[au + q(au)
ight]$$

The temperature at the depth point l is

$$T_l = \sqrt[4]{rac{3}{4} \cdot {T_{ ext{eff}}}^4 \cdot [au_l + q(au)]}$$

 $q(\tau)$ is called Hopf function.

The hydrostatic equation is solved in inside direction. The first four (outer) depth points are treated following the Runge-Kutta method. Then, the other points are calculated with a predictor-corrector method of Ham. In detail, the electron density at each depth point is calculated for the actual temperature and the given photospheric composition. The iteration method is based on Mihalas, Stellar Atmospheres, Second Edition, 5-2. Subsequently, LTE2 calculates LTE occupation numbers for all levels of all elements using the Saha-Boltzmann equation. From these values, the Rosseland mean opacity $\bar{\kappa}_R$ is determined.

$$rac{1}{ar{\kappa}_{
m R}} \equiv rac{\pi}{4\sigma_{
m R} T^3} \int_0^\infty \kappa_
u^{-1} \cdot rac{\partial B_
u}{\partial T} d
u$$

LTE2 gives as result the electron density as well as the total particle density for the whole atmosphere. The original $\log \tau$ scale is transformed into a $\log m$ scale (m is the mass column density, measured from the outer limit of the atmosphere).

In practice, the temperature stratification of the inner atmosphere differs evidently from a "real" LTE or NLTE stratification, even for pure continuum models without consideration of line blanketing. Thus, with these start models, PRO2 or LINE1 need a relatively large number of iterations to calculate the right temperature structure. In order to reduce this number of iterations, an Unsöld-Lucy temperature correction can be carried out until the temperature structure is stable. Some option can be given to control the temperature correction method. (Sect. 4.3.2).

4.3.2 The Options

With some directives in the input file DATEN, LTE2 can be controlled. Especially, the photospheric parameters line effective temperature $T_{\rm eff}$, surface gravity g, and photospheric abundances (which are necessary in any case) are selected.

The input file DATEN for the program LTE2 has the following structure (Sect. 5.4):

DATEN

 $T_{\parallel}EFF_{\parallel}T$

LOG_|G_|g

 $ABUNDANCE_{|}xx$ hh

T has to be given in Kelvin; g in cgs; xx element in TMAP code (Sect.2.1), hh abundance fraction by particle numbers.

Please note that $\sum_{i=1}^{\text{NATOMS}} hh_i = 1!$ LTE2 will check this. In the case that $\sum = 1$, LTE2 will adopt the given abundances unchanged; if not, LTE2 will re-normalize the given values! Please check STDOUT for respective information!

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TAU, SCALE, MINIMUM, -8.0*

This option sets the limit ($\log \tau_1$) of the outer atmosphere. This limit can be varied under the premise that the atmosphere has to be optically thin over the complete frequency grid at least at the first depth point. (This is one of the basic assumptions of PRO2 and LINE1. Attention: the requested value is only a start value which can be different from that of the finally calculated "LTE" model.

TAU, SCALE, MAXIMUM, 2.6*

This option sets the limit ($\log \tau_{\rm ND}$) of the inner atmosphere. This limit can be varied under the premise that the atmosphere has to be optically thick over the complete frequency grid at least at the inner depth point. (This is one of the basic assumptions of PRO2 and LINE1. Attention: the requested value is only a start value which can be differ from that of the finally calculated "LTE" model.

TAU SCALE x y z

This option defines the arrangement of the inner depth points.

- z = 0.0: depth points are logarithmically equidistant, $\Delta \tau = \frac{\tau_{\text{max}} \tau_{\text{min}}}{ND}$ (start values)
- z > 0.0: like z = 0.0, but the distance (start value) of the innermost depth points (89-90) is z (default is z = 1.0E 05)
- z < 0.0: like z = 0.0, but the distance of the depth points $(x \to y)$ decreases logarithmically by the factor z

ITMAX=100*

maximum number of iterations of the Unsöld-Lucy temperature correction method. In the case that a negative temperature is detected, LTE2 will stop immediately. The output model contains then the atmospheric structure of the iteration before.

DAMP=0.5*

start value of an artificially introduced damping factor within the Unsöld-Lucy temperature correction method. This factor is partly necessary because the method appears to be unstable in most cases. In the course of the iterations, this value is automatically reduced to increase the stability — this might seem to be convergence . . .

EPS=1.0E-6*

"convergence" (see above) limit (to stop the Unsöld-Lucy temperature correction)

LTE2_START_MODEL

This option is only valid if the file TIN is available which is the original file TOUT of a former LTE2 run and contains the temperature stratification of the previously calculated "LTE" model.

This option is helpful if a model has not reached the "convergence" limit (see above) in the first run and shall be brought to a better "convergence". Moreover, the "fake convergence" due to the automatically reduced damping factor can be checked.

OUTPUT, MODEL, FORMATTED

The output model is saved formatted.

```
PRO2_START_MODEL
```

This option is only valid if a PRO2 or LINE1 model is supplied as start model MIN. This is helpful to introduce new elements or to vary their abundance ratios. The temperature stratification of a convergent NLTE model is used here to calculate LTE occupation numbers (particle conservation) for all levels of all elements. With this option, ITMAX=0 is set automatically, to preserve the temperature structure. A follow-up line formation calculation (i.e. at fixed temperature) with PRO2 or LINE1 supplies a start model with NLTE occupation numbers and a — hopefully — good approximation of the final temperature stratification.

The following options can be given to control the output of LTE2. Their meaning should be clear ... XXXX can be substituted by EACH or LAST.

```
PRINT_BMERGENT_FLUX

PRINT_INTEGRATED_EDDINGTON_FLUX,ITERATION,XXXX

PRINT_OPTIONS

PRINT_MODEL_ATOMS_(OVERVIEW)

PRINT_LEVELS

PRINT_LTE_MODEL

PRINT_TEMPERATURE_CORRECTIONS,ITERATION,XXXX,ALL_DEPTHS

PRINT_TEMPERATURE_CORRECTIONS,ITERATION,XXXX,MAX.

PRINT_WARNINGS
```

The final model is saved by LTE2 into the file MODELL. The last record in MODELL is: MODPRG,DATE,TIME

(CHARACTER MODPRG*5,DATE*8,TIME*8 with MODPRG='LTE2', DATE and TIME with actual creation time) is added in order to identify the model. The temperature structure of the model is saved in the file TOUT (see above).

The program *LTE2* is available at Tübingen's PC/Workstation cluster in its latest version: /home/rauch/bimod/lte2.Linux_x64.

The actual PARAMETERs are:

```
• NATOMAX = 3

• NIONMAX = 9

• NLMAX = 139

• LTEMAX = 109

• NFMAX = 1299
```

Other executables of LTE2 with different PARAMETERs have to be created (Sect. 1).

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5 NLTE Models: PRO2

5.1 The Method

The program *PRO2* (Werner 1986; Werner et al. 2003; Rauch & Deetjen 2003; Werner et al. 2012) calculates plane parallel NLTE model atmospheres in radiative and hydrostatic equilibrium. It is based on the "Accelerated Lambda Iteration (ALI) method (Werner & Husfeld 1985). It consists mainly out of two iteration cycles. The "outer" cycle is called "Scharmer iteration" (this name is taken from Scharmer 1981 who worked on solutions of radiative transfer problems using approximate lambda operators) and iterates the radiation field according to

$$J^n_
u = \Lambda^* S^n_
u + (\Lambda - \Lambda^*) \cdot S^{n-1}_
u$$

The source function S_{ν} of the actual iteration n is depending on the radiation field J_{ν} which is to be calculated. To calculate S_{ν} , the non-linear statistical equations have to be solved under consideration of the radiative and hydrostatic equilibrium. This is done in the "inner" iteration cycle by a [quasi-] Newton-Raphson iteration. In the following this procedure is called "linearizations".

5.2 The PARAMETERs

For the compilation of PRO2 the files PARA.INC, PARA1.INC, and PARA3.INC have to be adjusted to the used atomic data file ATOMS (Sect. 2) and the frequency grid FGRID.

Example parameter files look like:

```
PARA. INC
    PARAMETER( NLMAX= 229,
                               NIONMAX= 79,
                                              NATOMAX= 17,
                 NFMAX=7999,
                               LTEMAX =931,
                                              NLATOM =229
                                                             )
 >
PARA1. INC
    PARAMETER (
                 NRBBMAX=249,
                                NRBFMAX= 229,
                                                NCBBMAX=599,
                                                               NCBFMAX=229,
                 NRFFMAX= 61,
                                NCBXMAX=4999,
                                                NRLLMAX=
                                                           1,
                                NRBFMAD=
                                                NCBBMAD=
 >
                 NRBBMAD=999,
                                            6,
                                                           6,
                                                               NCBFMAD=
 >
                 NRFFMAD=
                            2.
                                NCBXMAD=
                                            8,
                                                NRLLMAD=
                                                           1,
                                                               NRLUMAD=
                                                                              )
PARA3. INC
               NRDIMAX=1,
                            NRDIMAD=1,
                                         NSIG=599999 )
   PARAMETER(
```

The PARAMETER denote maximum numbers of e.g. elements etc. (see below) which can be treated with PRO2 if the files are used for its compilation, respectively.

parameter	maximum numbers of
NLMAX	NLTE levels
NIONMAX	ions
NATOMAX	elements
NFMAX	frequency points
LTEMAX	LTE levels
NLATOM	NLTE levels (block matrix method, for one element)
NRBBMAX	radiative bound-bound transitions
NRBFMAX	radiative bound-free transitions
NCBBMAX	collisional bound-bound transitions
NCBFMAX	collisional bound-free transitions
NRFFMAX	radiative free-free transitions
NCBXMAX	collisional bound-bound transitions (to LTE levels)
NRLLMAX	radiative transitions within a band of a complex ion
NRLUMAX	radiative transitions low→up of complex ions
NRBBMAD	data points in formula used for RBB
NRBFMAD	data points in formula used for RBF
NCBBMAD	data points in formula used for CBB
NCBFMAD	data points in formula used for CBF
NRFFMAD	data points in formula used for RFF
NCBXMAD	data points in formula used for CBX
NRLLMAD	data points in formula used for RLL
NRLUMAD	data points in formula used for RLU
NRDIMAX	radiative di-electronic transitions
NRDIMAD	data points in formula used for RDI
NSIG	size of array SIGBF (NRBF× frequency points)

NSIG uses the number of frequency points between the thresholds and the maximum energy. This reduces the size of he array SIGBF and saves core memory.

These PARAMETERS are given in the output of ATOMS2 and SETF2 (please use e.g. grep para to extract them for this output). The PARAMETER NSIG is named there NRBFMAW. In the PARAMETER files used for the compilation of LINE1, NSIG has to be eliminated from PARA3.INC and NRBFMAW has to be added in PARA1.INC accordingly.

5.3 Input and Output Files

The program PRO2 expects the following input files (only those which are marked with "*" are necessary):

$ATOMS^*$

atomic-data file (Sect. 2)

$DATEN^*$

contains all input cards, used as a here document for STDIN (file name can be freely chosen)

$FGRID^*$

frequency grid (Sect. 3)

$MODIN^*$

start model (Sect. 4)

RBF_CUTOFF

NLTE Models: PRO2

optional input file, that can be used to set specific bound-free absorption cross-section equal to 0, starting at a given frequency.

Example:

```
cat > RBF_CUTOFF << eos LOW___UP__CUTOFF-FREQUENCY C432S__UUUUU 1.0E16 HE28__HE31___1.2E17
```

The first record in *RBF_CUTOFF* (in the above example!) is optional.

The program PRO2 creates the following output files:

MODOUT

(output model)

NEGDENS

Indicates that negative density values exist in *MODOUT*. *MODOUT* can be read by *LINE1* and transformed into a formatted version. This model may be edited then.

STOP

Indicates that a model converged. PRO2 will stop immediately in case that this file exists in the working directory because it assumes convergence then (cf. Sect. 10). This can be used to stop PRO2 calculations (e.g. to restart the jobs with a changed input file DATEN.

5.4 Input Options

The program PRO2 can be controlled by a number of input options. Some of them are initialized with a *default* value (indicated by *) and can be omitted if no other values shall be requested. Indispensable blanks are indicated by ... Numerical data can be inserted *format free*.

. whatever

commented by "." in the first column, disregarded by PRO2

ABUNDANCE, AA x

introduces the abundance of the element AA (flushed to the left). The input x is the number ratio relative to hydrogen. If the start model contains already the element AA, this card is disregarded.

```
\mathtt{ABUNDANCE}_{\square}\mathtt{AA}_{\square}\mathtt{x}_{\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square}\mathtt{MASS}_{\square}\mathtt{FRACTION}
```

introduces the abundance of the element AA (flushed to the left). The input x is the mass fraction. If the start model contains already the element AA, this card is disregarded.

ACCELERATION,OF,CONVERGENCE,FROM,ITERATION 999*

the Ng method for acceleration of convergence is used, starting from iteration 999. After every 4th iteration, all photospheric parameters are extrapolated from the previous iterations.

This option invokes a quasi-Newton method ("Broyden method") instead of the Newton-Raphson iteration of the linearized equations.

- =0 Newton-Raphson iteration
- =1 Broyden method, at the beginning of each Scharmer iteration started by a normal Newton-Raphson iteration step. This is only suitable for diagonal operators ($\Lambda = 3$).
- =2 Broyden method, at the beginning of each Scharmer iteration started with the updated matrix of the last Scharmer iteration. This is only suitable for diagonal operators ($\Lambda = 3$).
- =3 like BROYDEN=1, but suitable for tridiagonal operators ($\Lambda = 4$).
- =4 like BROYDEN=2, but suitable for tridiagonal operators ($\Lambda = 4$).

The three switch-limits ϵ_1 , ϵ_2 , ϵ_3 indicate

- ϵ_1 switch from BROYDEN=0 \to 1, if the relative corrections of the Scharmer iteration are smaller than ϵ_1 . Is ($\epsilon_1 \ge 1.E33$) required, the 1st Scharmer iteration immediately starts with the Broyden method.
- ϵ_2 switch from BROYDEN=1 \rightarrow 2, if the relative corrections of the Scharmer iteration are smaller than ϵ_2 .
- ϵ_3 switch back to the next lower BROYDEN stage, if the relative corrections for the temperature or the electron density a higher than ϵ_3 . This shall avoid divergent iterations.

While in the case of diagonal operator the switches occur locally following local criteria, the tridiagonal operators require global judgement and used the highest relative correction found in the whole atmosphere. Informations about the switches can be printed to STDOUT (see below).

${\tt CHANGE_EFFECTIVE\ TEMPERATURE_x}$

generally, PRO2 takes the information about $T_{\rm eff}$ from the start model (Sect. 4). With this card, the another $T_{\rm eff}$ can be chosen. This works in small steps and takes much less time than the calculation of a new model with almost the same parameters.

$\mathtt{CHANGE}_{\square}\mathtt{LOGG}_{\square}\mathtt{x}$

like CHANGE EFFECTIVE TEMPERATURE, but for a new value of the surface gravity g

$\mathtt{CHANGE} _\mathtt{ABUNDANCE} _\mathtt{AA} _\mathtt{x} _ _ \mathtt{MASS} _\mathtt{FRACTION}$

like CHANGE EFFECTIVE TEMPERATURE, see ABUNDANCE for MASS FRACTION.

COMMENT:

With this card any comment can be printed at the beginning of the output.

DEPTH DEPENDENT LINE PROFILES, LINEARIZATION: XXXXX

This option XXXXX = FIRST or $EACH_{\sqcup}$ is indispensable if depth dependent line profiles shall be calculated. FIRST saves a lot of computational time.

NLTE Models: PRO2

ERRNEW=1.E-10*

limit for the relative corrections to stop the linearizations

ERRSCH=1.E-04*

limit for the relative corrections to stop the Scharmer iteration

FORMAL, SOLUTION

PRO2 carries out two formal solutions (with/without line opacities) and saves the line profile in the file LINES.

FREQUENCY GRID FORMATTED

The frequency grid FGRID (input) is formatted.

$$\mathtt{STEP}_{\square}\mathtt{UP}_{\square}\mathtt{F-VALUES}{:}_{\square}*****-\mathtt{START}{:}_{\square}\mathtt{IIII}_{\square}\mathtt{x}_1\ \mathtt{x}_2\ \mathtt{x}_3$$

The oscillator strengths of all lines of a selected ion IIII are reduced by a factor of x_1 at the beginning of the iterations and stepped up by a factor of x_2 every x_3 Scharmer iterations.

For **** the two possibilities INPUT or MODEL can be requested. With INPUT, PRO2 uses the values of x_1 , x_2 , and x_3 as requested in the card; with MODEL, PRO2 reads these values from the start model (Sect. 4) if a previous PRO2 has not reached unity for all step-up values and thus, written them to the output model.

STEP_UP_F-VALUES: NO KANTOROVICH RESET AFTER STEP_UP

In general, after each increase of f-values, the Broyden/Kantorovich method is reset. This card suppresses this reset.

$\mathtt{STEP}_{\square}\mathtt{UP}_{\square}\mathtt{F-VALUES}:_{\square}\mathtt{LIMIT}_{\square}\mathtt{FOR}_{\square}\mathtt{ABS}._{\square}\mathtt{REL}._{\square}\mathtt{NG}_{\square}\mathtt{CORRECTION}_{\square}\mathtt{x}$

In case that the absolute relative total density correction exceeds x (default value: 0.1), the next step up is skipped.

GAMMA=1*

The value defines the optical depth $\tau_{\nu} = \gamma$ which separates the region of the line wing for which the *core* and the *wing* approximations shall be used.

IGNORE, CPU, TIME, LIMIT

set the security time (necessary to complete the job) equal 0. This can only be used if it is sure that the requested number of iterations can be completed within the jobs run time limit.

${\tt INCREASE}_{\sqcup}{\tt COLLISIONAL}_{\sqcup}{\tt RATES}_{\sqcup}{\tt BY}_{\sqcup}{\tt FACTOR}_{\sqcup}{\tt x}$

The collisional rates can be increased in order to simulate a LTE model atmosphere. The factor x has to be $> 10^{+30}$, in order to guarantee that collisions dominate the photosphere . . .

INNER_BOUNDARY: LAMBDA-ITERATION

${\tt INPUT-MODEL}_{{\scriptscriptstyle |} \; {\scriptscriptstyle |}} {\tt FORMATTED}$

ITMAX=1*

maximum number of Scharmer iterations

JACOBI, FRESH-UP, INTERVAL 5*

calculation of the Jacobian every 5^{th} Scharmer iteration. Only valid in case of Broyden or Kantorovich method.

KANTOROVICH=2, SWITCH LIMITS 0-->1, 1-->2, 2-->1 : 1.E33* 0.1* 1.0* like the BROYDEN card.

- =0 Newton-Raphson iteration
- =1 Jacobian is calculated at the beginning of every Scharmer iteration but then kept constant.
- =2 Jacobian is calculated at the beginning of the 1st Scharmer iteration but then kept constant. A new calculation of the Jacobian is done following the JACOBI_□FRESH-UP card (see above)

LINEARIZE HYDROSTATIC EQUATION

With this option the hydrostatic equation is solved simultaneously with the statistical equation and not (default) solved subsequently.

NO RENORMALIZATION OF COMPLEX LINE CROSS SECTIONS

undoes a previously made re-normalization of the sample cross-sections for iron group elements

NO_TEMPERATURE_CORRECTION

the radiative equilibrium is omitted from the linearization

$$\texttt{T-CORRECTION}_{\square} \texttt{ONLY}_{\square} \texttt{IN}_{\square} \texttt{LOG}_{\square} \texttt{M}_{\square} \texttt{INTERVAL} \colon_{\square} m_{\min} \ m_{\max}$$

the temperature correction is restricted to the selected $\log m$ interval.

T-CORRECTION, PROFILE: X

the temperature correction is damped by simple functions (ND = total number of depth points)

 $\frac{X}{1}$ depth point / ND $\frac{1}{2}$ (depth point / ND)² $\frac{1}{3}$ $\sqrt{\frac{\text{depth point}}{ND}}$

LAMBDA=3*

selects the Λ operator

 $\Lambda = 3$ diagonal operator by Olson & Kunasz

 $\Lambda = 4$ tridiagonal operator by Olson & Kunasz

NLTE Models: PRO2

Both operators are generally parameter-free. In practice, it is less time consuming to do a Λ iteration in the optical thin case. This can be done by requesting a special GAMMA parameter (see above).

In order to accelerate the convergence, both operator switch to core saturation if $\tau_{\nu} > 100$.

LINEARIZATION, MODE: BLOCK-MATRIX-ITERATION, XX YY ZZ

with this option, only the occupation numbers of the given elements (here XX YY ZZ) are iterated. In combination with the card SOLVE STATISTICAL EQUATIONS ONLY, a "classical" line formation iteration is done. In all depth points, XX is Newton-Raphson iterated (inclusive constraint equations) first, then YY, and at least ZZ. Attention: this option alone does not switch off the constraint equations!

LINEARIZATION, MODE: BLOCK-MATRIX-ITERATION, ALL

like LINEARIZATION MODE: BLOCK-MATRIX-ITERATION, XX YY ZZ. All elements are iterated in order of their appearance in the atomic data file ATOMS.

MICROTURBULENCE, [KM/S] 0.0*

with this option, the microturbulence pressure is considered in the hydrostatic equation but **not** in the calculation of the line profiles!

NFWMAX=1*

maximum number of linearizations

${\tt OCCUPATION}_{\sqcup}{\tt PROBABILITY}_{\sqcup}{\tt FORMALISM}_{\sqcup}{\tt FOR}_{\sqcup}{\tt AA}$

The Hummer-Mihalas (HM) formalism is used to calculate the level dissolution for AA = H1 or $AA = \langle ion \rangle$.

OPACITY PROJECT RBF DATA START AT EDGE

PRO2 can calculate bound-free cross-section using data from the Opacity Project (OP). Here the OP data set is used and the cross-sections start at the threshold energy of the respective level.

${\tt OPACITY}_{\sqcup}{\tt PROJECT}_{\sqcup}{\tt RBF}_{\sqcup}{\tt DATA}:_{\sqcup}{\tt FULL}_{\sqcup}{\tt DATA}_{\sqcup}{\tt SET}$

see above, the complete OP data set is used (which may start at lower energy the the level energy in order to simulate the transition from the line absorption to the continuous absorption at the series limit.

$\mathtt{OPACITY}_{\square}\mathtt{PROJECT}_{\square}\mathtt{RBF}_{\square}\mathtt{DATA}:_{\square}\mathtt{ONLY}_{\square}\mathtt{EDGE}_{\square}\mathtt{VALUE}$

see above, a mean value of the OP data at the threshold energy is calculated, then the absorption cross-sections are calculated using the Seaton formula in hydrogenic approximation.

OPACITY, PROJECT, RBF, DATA: MISSING, HYDROGENIC

see above, for those levels which are not found in the OP data set hydrogen-like threshold cross-sections are calculated. Then, the Seaton formula is used.

OPACITY_PROJECT_RBF_DATA:_HYDROGENIC

see above, all bound-free transitions, which are found in the atomic data file ATOMS and request explicitly the use of OP data are calculated with hydrogen-like threshold cross-sections and the Seaton formula.

OUTPUT-MODEL, FORMATTED

RADIATIVE EQUILIBRIUM: DIFFERENTIAL/INTEGRAL FORM

The radiative equilibrium is calculated using a combination of the differential and the integral method.

READ, NEW, TEMPERATURE, STRUCTURE

A temperature structure is read in from file TEMPERATURE.IN. This temperature may be interpolated from other models in order to speed up the calculation of "in-between" models. TEMPERATURE.IN is an ASCII table with $\log m$ and T, however, the $\log m$ values are ignored.

$REDUCE_{||}LOG_{||}CVEC_{||}x$

This options scales the inhomogeneity vector of the linearized equations. This is useful to overcome numerical instabilities and to avoid too-large corrections in the beginning of the iterations. This card has to be inactivated to get a finally converged model.

SAVING MODEL: EACH ITERATION

a temporary model (MODTMP) is saved in the working directory \$TMPDIR. An example how to save this temporary model automatically in case of a failed model calculation is given in Sect. D.8.5. An

ln -sf MODTMP \${name}

before the start of the model calculation, where $\{name\}$ is the assigned name of the output model helps to identify the model.

SET TEMPERATURE t $\log m_1 \, \log m_2$ [scaling factor]

within $\log m_1 \leq \log m \leq \log m_2$, the model temperature is set to

t > 0: t

t = -1: cubic-spline interpolation of t

t = -2: linear interpolation of t

t = -3: t read from model TIN

t=-4: t is temperature value of next depth point with $\log m > \log m_2$.

t = -5: like t = -3, but an additional third argument (scaling factor) has to be given that is multiplied to the TIN temperature

SHIFT EDGE BBBBBBBBBBFFFFFFF x

The absorption threshold of the level BBBBBBBBBFFFFFFFF given in the TMAP code (Sect. 2.1) is artificially shifted to the frequency point x. Attention: The given frequency x has to fit exactly a point within the frequency grid FGRID! All shifts are printed to STDOUT.

NLTE Models: PRO2

SKIP ATOM AA BELOW x

the element AA (flushed to the left) with occupation numbers less than x are eliminated from the statistical equations

SKIP, ION, III, BELOW x

ion III (flushed to the left) with occupation numbers less than x are eliminated from the statistical equations

SKIP LEVELS BELOW x

levels with occupation numbers less than x are eliminated from the statistical equations

SKIP, OCCDRVF, AFTER, 1ST, LINEARIZATION

The derivations of the source functions are only calculated once (in the 1st linearization) and are then kept fixed. Attention: This saves an enormous amount of computational time in case of many NLTE levels and many frequency points but it works well only in the case of almost converged models.

SOLVE, STATISTICAL, EQUATIONS, ONLY

switches off the constraint equations for n_e, n_H, n_g, T

SOLVE STATISTICAL EQUATIONS ONLY RE-SOLVE PARTICLE CONSERVATION

switches off the constraint equations for n_e , n_H , n_g , T within the Newton-Raphson iteration. The hydrostatic and particle conservation equations are then solved subsequently.

SWITCH OFF LINES

All line transitions are ignored. Attention: sample lines of iron-group elements can not be switched off.

TIME, LIMIT 2000*

cpu time for the job to calculate. Only valid on non-CRAY machines

UNSOELD-LUCY TEMPERATURE CORRECTION DAMP x

With this option, the radiative equilibrium is eliminated from the linearization. The temperature structure is calculated after the linearizations with the Unsöld-Lucy temperature correction method. The damping factor x is used to avoid over-corrections and numerical instabilities.

$\texttt{UPPER}_{\square} \texttt{AND}_{\square} \texttt{LOWER}_{\square} \texttt{LIMITS}_{\square} \texttt{FOR}_{\square} \texttt{RELATIVE}_{\square} \texttt{T-CORRECTION}_{\square} \texttt{t}_1 \enspace \texttt{t}_2$

The relative temperature corrections (iteration i) are limited to the range $T_i \cdot t_1 \leq T_{i+1} \leq T_i \cdot t_2$ at

all depth points.

5.5 Output Options

The following PRINT and PLOT cards create output on STDOUTt and plot data files, respectively. The plot files are written in WRPLOT readable format. The option should be easily understandable ... Generally, for XXXX EACH or LAST can be inserted. ii indicates an I2 format specifier (FORTRAN).

5.5.1 Output to STDOUT

PRINT, ABUNDANCES

PRINT, BROYDEN, INFORMATIONS

PRINT_CORRECTIONS_OF_LAST_LINEARIZATION, ITERATION:LAST, DEPTH_INCREMENT: i

PRINT CORRECTIONS OF TOTAL DENSITIES, ITERATION: LAST, DEPTH INCREMENT: i in the case of non-linearization of the hydrostatic equation (default, see above)

PRINT, CP-TIME/ITERATION, XXXX

 $\mathtt{PRINT}_{\square}\mathtt{DATA}_{\square}\mathtt{FOR}_{\square}\mathtt{DIELECTRONIC}_{\square}\mathtt{RECOMBINATIONS}$

prints especially the frequencies which are selected by PRO2.

 $\mathtt{PRINT}_{\square} \mathtt{DEPARTURE}_{\square} \mathtt{COEFFICIENTS}, \mathtt{ITERATION:XXXX}, \mathtt{DEPTH}_{\square} \mathtt{INCREMENT:15}$

 $\mathtt{PRINT}_{\sqcup}\mathtt{EMERGENT}_{\sqcup}\mathtt{FLUX}\,\mathtt{,}\,\mathtt{ITERATION}\,\mathtt{:}\,\mathtt{XXXX}$

prints also T_{eff} , which is an sensitive control parameter

 ${\tt PRINT}_{\sqcup}{\tt FREQUENCY}_{\sqcup}{\tt GRID}$

 $\mathtt{PRINT}_{\square}\mathtt{INFORMATION}_{\square}\mathtt{ABOUT}_{\square}\mathtt{INPUT}_{\square}\mathtt{SAMPLE}_{\square}\mathtt{CROSS}_{\square}\mathtt{SECTIONS}$

 ${\tt PRINT}_{\sqcup} {\tt INPUT}_{\sqcup} {\tt MODEL}, {\tt DEPTH}_{\sqcup} {\tt INCREMENT} \colon {\tt ii}$

 $\texttt{PRINT}_{\square} \texttt{INPUT}_{\square} \texttt{MODEL}, \texttt{DEPTH}_{\square} \texttt{INCREMENT} : \texttt{ii}_{\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square} (\texttt{STRUCTURE}_{\square} \texttt{ONLY})$

PRINT_INTEGRATED_SURFACE_FLUX, ITERATION: XXXX

PRINT, KANTOROVICH, INFORMATIONS

PRINT, LEVELS

prints level names, energies, and statistical weights of all LTE / NLTE levels

PRINT LEVEL SKIP INFORMATION

PRINT_MAX._REL._CORRECTIONS_EVERY_ii_ITERATIONS

PRINT, OCCUPATION, PROBABILITIES, ITERATION: XXXX, DEPTH, INCREMENT: ii

PRINT, OPACITY, PROJECT, INFORMATION

PRINT, OUTPUT, MODEL, ITERATION: XXXX, DEPTH, INCREMENT: ii

prints temperature and density structure, and all occupation numbers of the NLTE levels

 $\mathtt{PRINT}_{\sqcup}\mathtt{OUTPUT}_{\sqcup}\mathtt{MODEL}\mathtt{,ITERATION:XXXX}\mathtt{,DEPTH}_{\sqcup}\mathtt{INCREMENT:ii}_{\sqcup\sqcup\sqcup}\mathtt{(STRUCTURE}_{\sqcup}\mathtt{ONLY}\mathtt{)}$

prints temperature and density structure

PRINT, MODEL, ATOMS, (OVERVIEW)

PRINT NUMBER OF FUV PHOTONS

PRINT, OPTIONS

prints the most important input parameters

PRINT PARAMETER NSIG

for the optimization of the PARAMETER NSIG, NRLUMAD, and NRLLMAD

 $\mathtt{PRINT}_{\square}\mathtt{RADIATION}_{\square}\mathtt{FIELD}, \mathtt{ITERATION}:\mathtt{XXXX}, \mathtt{DEPTH}_{\square}\mathtt{INCREMENT}:\mathtt{ii}$

PRINT, RBF, CUTOFF, FREQUENCIES

prints the frequency where the RBF cross-section of a level is set equal 0

 ${\tt PRINT_{\tiny ||}} {\tt ROSS, ITERATION: LAST}$

print the value of ROSS for the inner boundary condition

PRINT, TAUSCALES, ITERATION: XXX, DEPTH, INCREMENT: ii

PRINT, WARNINGS

LP-PLOT OPTICALLY THICK/THIN, ITERATION: XXXX

A plot of the $\tau = 1$ limit in the atmosphere is printed and saved in a file (*PLLP*).

5.5.2 Output into Plot Data Files

PLOT, CORRECTIONS

 ${\tt PLOT_{\scriptscriptstyle | \, |}} {\tt DEPARTURES} \ {\tt xmin} \ {\tt xmax} \ {\tt ymin} \ {\tt ymax}$

 $\label{eq:plot_loss} PLOT_{\sqcup\sqcup} \\ EMERGENT_{\sqcup} \\ FLUX, ITERATION: XXXX \\ \\ same like PRINT card, output is written into file PRFLUX \\$

 $PLOT_{,,}FLUX$ xmin xmax ymin ymax

PLOT IONIZATION FRACTIONS AA xmin xmax ymin ymax

$$\label{eq:locally_local} \begin{split} & \texttt{LP-PLOT}_{\square} \texttt{OPTICALLY}_{\square} \texttt{THICK/THIN,ITERATION:XXXX,FILE}_{\square} \texttt{ONLY} \\ & \texttt{A plot of the } \tau = 1 \text{ limit in the atmosphere is saved in the file } \textit{PLLP} \text{ only.} \end{split}$$

PLOT TEMPERATURE STRATIFICATION

CUTM and NDSCALE 37

6 Adjust the log m Intervals of Existing Models: CUTM and NDSCALE

6.1 Adjust $\log m$ Intervals of Models

One boundary condition of TMAP is that at least the outer two depth points have to be optically thin and at least the inner two depth points have to be optically thick.

For the calculation of model grids, it is an advantage to use approximately the same $\log m$ interval for all models as well as the frequency grids (see above).

In case of numerical instabilities it may be helpful to adjust the $\log m$ interval of existing models. This is done by CUTM (to reduce the $\log m$ interval, both minimum and maximum) and EXTEND (to reduce the minimum $\log m$).

Since the number of depth points will deviate from the standard of 90, NDSCALE can interpolate models with any number of depth points to a new models with a chosen number of depth points.

6.1.1 Reduce the Depth Interval: CUTM

First copy a binary TMAP model (90 depth points) to the file MODIN, and start e.g. /home-/rauch/bimod/cutm.Linux_X86. CUTM asks for the log m interval and writes the reduced model to the file MODOUT. The following displays an example batch-job file for CUTM:

```
#!/bin/sh
          ${HOME}/.jobstart
set +x;.
# do not edit the beginning of this file
##########################
## own job following ##
########################
cp <binary input model> MODIN
/home/rauch/bimod/cutm${sys} << eos
-7.50
+2.00
eos
mv MODOUT <binary output model>
# do not edit the rest of this file
#####################
set +x; ${HOME}/.jobend ${TMPDIR}
```

6.1.2 Extend a Model Atmosphere to the Outside: EXTEND

Start with the creation of a formatted TMAP model and copy it to the file MODIN. Write the number ND of depth points of this models into the file $MODIN_ND$. and start e.g. /home-/rauch/bimod/extend.Linux_X86. EXTEND asks for the log m minimum and writes the extended model to the file MODOUT. The following displays an example batch-job file for EXTEND:

```
#!/bin/sh
   set +x;.
             ${HOME}/.jobstart
  # do not edit the beginning of this file
   ##############################
   ## own job following ##
   #########################
   echo ''90'' > MODIN_ND
   cp <formatted input model> MODIN
   /home/rauch/bimod/extend${sys} << eos
   -8.00
   eos
  mv MODOUT <binary output model>
   # do not edit the rest of this file
   #####################
   set +x; ${HOME}/.jobend ${TMPDIR}
```

6.1.3 Interpolate Models for a New Depth-Point Number: NDSCALE

NDSCALE can interpolate a model to a new number of depth points. The may be necessary in case that the $\log m$ interval of a model was adjusted (Sect. 6.1).

Copy a binary TMAP model to the file MODIN, and start e.g. /home/rauch/bimod/nd-scale.Linux_X86. NDSCALE asks for the old and new number of depth points and writes the interpolated model to the file MODOUT. The following displays an example batch-job file for NDSCALE:

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7 Interpolate between Existing Models: INTER

7.1 Interpolate between Existing Models

/home/rauch/bimod/inter.Linux_<X86|P4|x64>. *INTER* read two models, *MOD.1* and *MOD.2*, that have been calculated with the same ATOMS and same number of depth points, and linearly interpolates them to MOD.I. This model can be used ideally as a start model for PRO2.

8 NLTE Line Formation and Models: LINE1

The program *LINE1* calculates — based on a suitable start model (Sect. 3) — model atmospheres like *PRO2*, NLTE occupation numbers at fixed temperature and (optional) density stratification (e.g. for new levels in the case of extended model ions), and theoretical line profiles of selected lines under consideration of the most recent line broadening theories.

The input and output options are generally the same like those of PRO2 (Sect. 5.4 and 5.5).

8.1 Input Files

The program *LINE1* expects the following input files (only those which are marked with "*" are necessary):

ABUNDi

It is possible to read occupation numbers for newly implemented levels from 14 additional, already existing models ($\rightarrow MODINi$). i (from ABUNDi) is a number from 1 to 9 (model 1 is MODIN, with abundance file **ABUND**, model 2 is MODIN2, with abundance file **ABUND2**, etc.) and 0, A, B, C, D, E for 10, 11, 12, 13, 14, and 15, respectively. Abundance ratios are taken from the first model that includes the element, i.e., with ascending priority from 1^{st} , 2^{nd} , ..., 15^{th} model. If the elemental abundances of the input model ($\rightarrow MODIN$) and of the models MODINi differ, scaling factors can be given in the files ABUNDi which are applied to the occupation numbers of the model MODINi, respectively.

Example:

 $C_{1}2$

 $\mathbb{N}_{\sqcup}^{\square} 1.3$ It is recommended to add the complete set of occupations numbers to a model that does not include the respective species. Otherwise the lower levels may artificially be underoccupied.

ADJUST

In practice NLTE model atmosphere calculations have shown that is is not possible in all cases to consider all line transitions simultaneously from the beginning. One method to overcome this problem is to reduce the oscillator strengths at the beginning with reduction factors which increase to unity in the course of the iteration.

The input file *ADJUST* contains the reduction factors and the factors with which the reduction factors are increased every iteration until the original oscillator strengths are reached.

Example:

means:

the oscillator strengths of all C IV line transitions are reduced by a factor of 10^{-5} . This factor is constant over the whole calculation. Additionally, the C IV 2s-2p line is individually reduced by the factor of 0.01 (\rightarrow total of 10^{-7}) but this factor (0.01) is increased by 5% every iteration. The C IV 5s-5p line is considered with its correct oscillator strength ($10^{-5} \cdot 10^{+5} = 1$). In the case that the temperature correction yields negative temperatures, the oscillator strengths are kept at their last value.

ATOMS*

atomic data file (Sect. 2), like for the start model or extended ...

DATEN

input and output options (Sect. 5.4, 5.5, and 8.3)

this files has to be copied to stdin because LINE1 reads the options from there (e.g. /home/rauch/bimod/line1 < DATEN).

DISTA

The frequency grid for the detailed calculation of line profiles is created by *LINE1* itself. The discretization of the frequency points within the line transitions can be split in intervals with different distances from point to point in order to represent the line centers or possible forbidden components (e.g. \rightarrow He I $\lambda 4471\text{Å}$) with a sufficient number of (narrowly spaced) frequency points. *DISTA* contains informations how many points at which distance ($\Delta\lambda$ in Å) shall symmetrically (to the line center) be inserted in these intervals.

Example:

5 0.1

10 0.3

15 0.5

results in points at $(\Delta \lambda \text{ in Å, only the "red" wing given})$:

0.0, 0.1, 0.2, 0.3, 0.4, 0.7, 1.0, 1.3, 1.6, 1.9, 2.4, 2.9, 3.4, 3.9, 4.4

In case of many lines and a high resolution, the parameter NRBBMAD (Sect. 1) may become very high. Due to compiler limitations in the size of arrays, no executable may be created then. To avoid this, the use of a respective F_BASE file (Sect. 3) and an adjusted DISTA is recommended. Example:

 F_BASE creates an equidistant frequency grid from 3000 Å to 7000 Å ($\Delta\lambda = 0.1$ Å), then a DISTA

2 0.01

4 0.02

6 0.03

8 0.05

9 100.00

10 0.01

is sufficient (narrow points only until the F_BASE resolution is reached, then just an end point of the line and a CONT/(BLUE/RED) point (Sect. 3). In addition, a dynamical use of DISTA in $LINE1_PROF$ is helpful and is described in Sect. 9.

$FGRID^*$

frequency grid (Sect. 3), not necessary in the case of line profile calculations (see above)

ION

contains the code for the selected ion (line profile calculations). It differs from the TMAP code (Sect. 2.1): the element is given in CHARACTER*2 format (flushed to the left) and the ionization stage is indicated in the third (and fourth) column.

Examples:

HE2

 $C_{1}2$

CA10

SETF2 creates this file for a given atomic-data file (Sect. 3).

LINIEN this file contains information about the range which has to be considered in the calculation of the line profiles and the selected line to calculate:

BLENDRANGE 40.0*

This card is of special importance and has to be the first record in LINIEN.

- LINE1 searches in the interval $[\lambda_o \lambda_{Blendrange}, \lambda_o + \lambda_{Blendrange}]$ for line frequency points (Sect. 3) of other line transitions. All line transitions with points between CONT RED and CONT BLUE of the selected line (see below), are regarded as a blend and included in the calculation in detail. All other line transitions in the input file ATOMS (see above) are disregarded in the creation of the internal atomic data and frequency grid files by LINE1. This minimizes both, core memory and computational time requirements.
- LINE1 considers in the creation of the internal atomic data and frequency grid files only those radiative bound-free (RBF) transitions with an ionization energy smaller than $c/(\lambda_o \lambda_{Blendrange})$ because RBF transitions with a higher ionization energy do not contribute to the opacity within the blendrange. This minimizes also the core memory and computational time requirements.

XXXXXXXXXYYYYYYYYYY

TMAP code (Sect. 2.1) of selected lines for which theoretical line profiles shall be calculated SETF2 creates this file for a given atomic-data file (Sect. 3).

MODIN*

start model (Sect. 4)

MODINi

see ABUNDi

OP_RBF_XXXX

bound-free cross-sections from the Opacity Project for levels of the ion XXXX

STARK_H1

VCS-like Stark broadening tables for H I

H_I line-broadening has changed in since 2008. The reason is that an error (for high members of the spectral series only) in the H_I line-broadening tables by Lemke (1997) that were used before. Tremblay & Bergeron (2009) and Tremblay & Bergeron (2015, priv. comm.) provide, parameter-free Stark line-broadening tables for H_I considering non-ideal effects. These replaced Lemke's data for the lowest members of the Lyman (n - n' = 1 - [2 - 21]), Balmer (n - n' = 2 - [3 - 22]), Paschen (n - n' = 3 - [4 - 22]), and Brackett (n - n' = 4 - [5 - 14]) series. These profiles can only consistently used with the Hummer & Mihalas (1988) equation-of-state (cf.,

OCCUPATION PROBABILITY FORMALISM FOR H1

in Sect. 5.4). For higher series members, we use the Holtsmark approximation.

STARK_ALIII

tables for He I (Dimitrijević & Sahal-Bréchot 1993a)

STARK_HE1

tables of Barnard et al. (1969) and Griem (1974) for He I

STARK_HE2

VCS-like tables for He II (Schöning & Butler 1989a,b)

STARK_C4

tables for C IV (Schöning 1993)

STARK_N5

tables for N v (Schöning 1995)

STARK_HEI

tables for He I (Dimitrijević & Sahal-Bréchot 1990)

STARK CIV

tables for C IV (Dimitrijević et al. 1991b; Dimitrijević & Sahal-Bréchot 1992c)

STARK_CV

tables for C v (Dimitrijević & Sahal-Bréchot 1996)

STARK NV

tables for N v (Dimitrijević & Sahal-Bréchot 1992b)

STARK_OIV

tables for O IV (Dimitrijević & Sahal-Bréchot 1995)

STARK_OV

tables for O v (Dimitrijević & Sahal-Bréchot 1995)

STARK_OVI

tables for O vi (Dimitrijević & Sahal-Bréchot 1992a)

STARK_SIIV

tables for Si IV (Dimitrijević et al. 1991a)

STARK_SVI

tables for S vi (Dimitrijević & Sahal-Bréchot 1993b)

Note: For H I, He II, C IV (STARK_C4), and N V STARK_N5 the tables have only to be loaded. *LINE1* uses these data files if they are available. For the other ions an option has to be given (see below) in *DATEN* to use the broadening tables for all lines of the ion, or for selected line transitions by requesting formula 5 (keyword RBB) in *ATOMS* (Sect. 2).

8.2 Output Files

The program LINE1 creates (besides STDOUT) the following plot data files (in accordance with the options given). Note that in general wavelengths $\leq 3000\,\text{Å}$ are given as vacuum wavelengths and wavelengths $> 3000\,\text{Å}$ as air wavelengths.

ANGJ_C.DAT

specific intensity (continuum only) for different angles (see PARAMETER NA, Sect. 1)

ANGJ_L.DAT

specific intensity for different angles (see PARAMETER NA, Sect. 1)

PLDEP

departure coefficients for selected levels

PLDIFF

abundance profiles for all elements

PLEFL

emergent flux (complete frequency interval)

PLLP

limit $\tau = 1$ (complete frequency interval)

PLPRF

line profiles

PLRBF

bound-free cross-sections of selected levels

PLWF

emergent flux $(F_{\lambda}(\lambda) - WF \text{ denotes wavelength} + \text{flux}.$

PLWFP

emergent flux $(F_{\lambda}(\lambda))$ and relative flux $F_{\lambda}/F_{\text{cont}}$ – WFP denotes wavelength + flux + normalized flux (profiles ...).

PLWFP_VACUUM

emergent flux $(F_{\lambda}(\lambda))$ and the relative flux $F_{\lambda}/F_{\text{cont}}$ – WFP_VACUUM denotes wavelength + flux + normalized flux (profiles ...). All wavelengths are vacuum wavelengths.

PLWP

emergent flux $(F_{\lambda}(\lambda) - \text{WP denotes wavelength} + \text{normalized flux (profiles ...)}).$

STRUCTURE

temperature stratification

For the evaluation and visualization of the output and plot data files several auxiliary programs are available (Sect. 11).

8.3 Options

For *LINE1* most of the options described in Sect. 5.4 and 5.5 are valid. There are some further options. Their meaning should be clear . . .

${\tt ACCEPT_CHANGED_EFFECTIVE_TEMPERATURE_X}$

LINE1 generally takes $T_{\rm eff}$ from the start model (MODIN). For very large models the change of $T_{\rm eff}$ (in small steps) takes much less times than the complete new calculation with mainly the same (!) parameters. However, still some iterations are needed . . .

${\tt ACCEPT}_{\sqcup}{\tt CHANGED}_{\sqcup}{\tt ABUNDANCES}$

LINE1 generally takes the chemical composition from the start model (MODIN) and ignores ABUNDANCE cards in the input file but for newly implemented elements. For very large models the change of the abundances (in small steps) takes much less times than the complete new calculation with mainly the same (!) parameters. However, also still some iterations are needed . . .

CPU-TIME, TTTTT

maximum CPU time for the job, TTTTT given in seconds $\mathtt{CPU-TIME}_{_}0.0$

This card transfers the total cpu time (then: stop) to LINE1. There are load dependent variations

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of the system cpu time which accounts to the user cpu time. These variations can effect the security time needed by *LINE1* in order to complete the iteration, write the model, etc.

DOPPLER, PROFILES

This option sets the line broadening of all line transitions in the atomic data file ATOMS generally to Doppler line broadening. This can be used for a quick overview about blends within a selected range.

FORCE USE OF LINE BROADENING TABLES

This option enables the use of line broadening tables instead of approximate formulae during the model atmosphere calculation.

ITERATION IN DEPTHS ii jj kk

The the first kk iterations are carried out only between depth points ii – jj.

LINE, TEMPERATURE, FROM, FORMATION, DEPTH

The line temperature T_{line} is by defaults set to $\frac{3}{4} \cdot T_{\text{eff}}$ (Sect. 3). The different lines form in very different depths, i.e. at different temperatures. This card allows to select the temperature at the formation depth ($\tau = 1$) of the line core for every lines as line temperature, respectively.

LINEARIZATION, MODE: BLOCK-MATRIX, ITERATION, ION

analogously to LINERIZATION $_{\!\!\!\perp}$ MODE ... in Sect. 5.4. This card allows to select one single ion to iterate.

MATRIX, INVERSION: MINV

The matrix inversion for the solution of the statistical equations is carried out by the SCILIB routine MINV. Attention: numerically instable for $n > 200, \ldots, 220$.

MATRIX, INVERSION: INV

The matrix inversion is done by an own routine.

MATRIX, INVERSION: GIRL

The matrix inversion is done by routine GIRL (single precision). More stable than MINV in some cases!

MATRIX, INVERSION: GIRLDP

The matrix inversion is done by routine GIRL (double precision).

MATRIX, INVERSION: SGETRF, + SGETRI

The matrix inversion is done by the LAPACK routines SGETRF and SGETRI.

MISSING OCCUPATION NUMBERS FROM I MODELS

The occupation numbers of newly considered levels are read from I models ($I = \max i$, see ABUNDi and MODINi, above) in increasing order. A consequence of NLTE modeling is the over-population of low-lying levels. A strong extension of models towards higher levels will not consider their de-population and may underestimate the population of the lowest. In this case, use the REPLACE, OCCUPATION, NUMBERS card (see below).

MIKROTURBULENCE [KM/SEC] 0.0

analogously to Sect. 5.4. *LINE1* considers the microturbulence also in the calculation of the line profiles.

NEWMAXB= 20*

maximum number of linearizations of the Broyden or Kantorovich iterations

REPLACE, OCCUPATION, NUMBERS, FOR, AAAA, FROM, MODEL, I

Occupation numbers of model I will be used for all levels of ion AAAA.

USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_C_IV

With this option, all C IV line transitions in the atomic data file *ATOMS* are calculated with the stark broadening tables by Dimitrijević (if found there).

If the Dimitrijević tables shall be used only for selected lines, in *ATOMS* formula 5 instead of formula 3 or 4 has to be requested. (Sect. 3).

— The tables (Sect. 8.1) have to be loaded in any case ...—

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_AL_III$

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_C_V$

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_HE_I$

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_N_V$

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_O_IV$

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_O_V$

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_O_VI$

 $\tt USE_DIMITRIJEVIC_BROADENING_TABLES_FOR_P_V$

USE DIMITRIJEVIC BROADENING TABLES FOR SI IV

USE, DIMITRIJEVIC, BROADENING, TABLES, FOR, S, VI

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all these cards like $\mathtt{USE}_{\square}\mathtt{DIMITRIJEVIC}_{\square}\mathtt{BROADENING}_{\square}\mathtt{TABLES}_{\square}\mathtt{FOR}_{\square}\mathtt{C}_{\square}\mathtt{IV}.$

PRINT, ATOMIC, DATA, FILE, ATOMS_2

print the internally created atomic data file (see above) for the line profile calculations

PRINT, EQUIVALENT, WIDTH

A small table with the equivalent widths of all selected lines is printed to STDOUT.

Attention: For the calculation of the equivalent widths of the selected lines, the interval [CONT RED,CONT BLUE] (Sect. 3) is used — with all included blends In order to study the equivalent widths of selected lines in detail the blendrange has to be set to a small-enough value that includes only the components of the multiplet or the atomic data file ATOMS has to be created with only the requested line transition.

PRINT, EQUIVALENT, WIDTH, (LONG)

like PRINT_EQUIVALENT_WIDTH but this table includes all frequency points in the interval [CONT RED,CONT BLUE] and gives additionally the accumulated equivalent widths.

PRINT, MULTIPLET, SPLITTING

LINE1 is able to split the NLTE occupation number of multiplets (see above). With this card all multiplet splittings are reported.

PRINT, VCS, TABLE

PRINT_FORMATION_DEPTH_OF_LINES_AND_THRESHOLDS

$\mathtt{PRINT}_{\square}\mathtt{FREQUENCY}_{\square}\mathtt{GRID}_{\square}\mathtt{FGRID}_{-2}$

print the internally created frequency grid (see above) for the line profile calculations

PRINT, LEVELS, WITH, LTE, START, VALUES

PRINT_LINES_AND_BLENDS

PRINT, PROFILE, TYPES

With this option, information about the used line-broadening theories for all calculated lines is printed to STDOUT. The same information is always included in the plot data file *PLPRF* as far as the necessary plot option (see below) is active to create it.

 $\mathtt{PRINT}_{\square}\mathtt{STATISTICS}_{\square}\mathtt{OF}_{\square}\mathtt{FGRID}_2$

PRINT, WARNINGS

PLOT_EMERGENT_FLUX__LOG_FNUE__/_LOG_NUE

PLOT_EMERGENT_FLUX__LFNUE__/_NUE

PLOT_EMERGENT_FLUX__LOG_FLAM__/_LOG_LAM

PLOT_EMERGENT_FLUX__FLAM__/_LAM

PLOT_EMERGENT_FLUX__FLAM__/_LAM__INTERVAL: LAMBDAMIN LAMBDAMAX

PLOT, LIMIT, TAU=1, ITERATION: EACH

With this option, a plot file PLLP is created which contains the geometrical depth (logarithmical) of the limit $\tau = 1$ for the complete frequency grid. It can be plotted with the program PLXY (Sect. 11).

PLOT_LINES

 $\label{eq:plot_temperature_stratification, iteration: last \\ PLOT_temperature_stratification, iteration: each \\$

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9 Synthetic Spectra: LINE1_PROF

The program LINE1_PROF calculates — based on a suitable start model (Sect. 3) — a synthetic spectrum and/or line profiles of selected lines under consideration of the most recent line-broadening theories.

The input and output options are the same like those of LINE1 (Sect. 8).

DIFFUSION MODEL

read a diffusion model calculated by NGRT.

In general, an atomic data file for a selected wavelength range (in order to limit the number of frequency points) should be created using ATOMS2, e.g. with the card:

The use of IRONIC data for iron-group elements requires to use the same frequency grid for the calculation of the synthetic spectrum. Thus,

PRINT, ENTSTEHUNGSTIEFEN, DER, LINIEN, UND, KANTEN

LINE1_PROF prints the formation depths of line cores and ionization edges

PLOT, IONIZATION, FRACTIONS

LINE1_PROF writes ionizations fractions of all ionic species in ATOMS to the file PLION that may be saved to e.g.<name>. Using

```
ln -sf <name> ion2devide.dat
/home/rauch/bin/ion2devide.Linux_X86
```

yields a series of ionfrac.<II> (II is the ionic species). These files can be plotted together in one plot with the template

/home/rauch/tools/IonFrac.wr,

that has to be edited properly for the individual star.

SAVE, FGRID_2

allows to save the frequency grid created by $LINE1_PROF$ is saved into the files $FGRID_2$ (binary) and $FGRID_3$ (ASCII). $FGRID_3$ can be used by IRONIC then. In order to check the respective ATOMS file, SETF2 has to be used to create a "start" input FGRID file.

 $LINE1_PROF$ considers the same input files like SETF2, $CONTS_MAN$, F_BASE , and POS_LIST .

A difference is the content of F_BASE , where

 λ_{\min} λ_{\max} $\Delta\lambda$

have to be given. There may be entries for different intervals.

LINE1_PROF uses an external frequency grid, if

FGRID_2, FROM, INPUT

is activated.

 ${\tt DYNAMIC_DISTA} \ \lambda_{\min} \ \lambda_{\max} \ {\tt DYNFACTOR}$

In general, spectral lines are more narrow in the X-ray or UV compared to the IR. To avoid excessively high NRBBMAD values, a dynamic adjustment of the blue and red line limits is necessary. The values given in the input file DISTA are inter/extrapolated to DISTA values $\times 1$ at λ_{min} and DISTA values $\times DYNFACTOR$ at λ_{max} .

The program *LINE1_PROF* creates (besides STDOUT and the files mentioned in Sect. 8) the following plot data files (in accordance with the options given).

IDENT

for each line within the BLENDRANGE (Sect. 8), an

\IDENT <wavlength> <ion> <transition>

card is created that can be used by WRPLOT (Sect. 11) for line identifications.

To combine subsequent IDENTs with identical ion name, use /home/rauch/tools/mkID-MULT.Linux_p64 with an input file id.in. The results will be in an output file id.out.

PLDIFF

element mass fractions in case of a diffusion input model

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10 Clean Up

TMAP requires a controlled clean up from time to time.

10.1 Delete TMAP Working Directories

The general job structure of TMAP is

```
# do not edit the rest of this file
########################
set +x; ${HOME}/.jobend ${TMPDIR}
```

```
$\{\text{HOME}}/.jobstart creates a working directory of the form /scratch/weekly/<username>/YYYY-MM-DD_hh:mm:ss that is deleted at the end of a job by $\{\text{HOME}}/.jobend. In case of a user break (<CTRL C>), the working directory and all data therein remain on the scratch disk. These have to be deleted regularly by
```

 \sim rauch/tools/clearTMAPscratch@all_linux_no_ask .

Be aware, that this command deletes all TMAP working directories. In case that TMAP jobs are still running, they will fail!

10.2 Kill TMAP Jobs

In case that TMAP jobs were started using the nohup command, they will continue even if the user logged out or if the console was rebooted. Then, these jobs can be killed individually on the respective machines (cf. user of the STOP file, Sect.. 5), or with the procedure

~rauch/tools/kill@all_linux_no_ask .

Be aware, that this command kills all your jobs (not only TMAP jobs) on all linux machines but at your local machine!

11 Auxiliaries

Three plot programs, PROF1, and PLXY, are available on the computers at Tübingen. They are designed for the visualization of plot data files from the Tübingen NLTE Model-Atmosphere Package (TMAP). The program PLXY is able to plot any x-y table. The additional program WR-PLOT is a plot program which is not exclusively based on the Tübingen NLTE Model-Atmosphere Package (TMAP). The two auxiliary programs RTFZ and PLADD are designed to interactively rectify and co-add spectra.

The programs are to be used interactively and allow an individual design of the plot. Interactive labeling etc. is possible. All interactively inserted label can be saved in files program.AUX and can be re-loaded if the program is invoked again later.

The plot files are created in PostScript format, EPSF is optional for a later inclusion in \LaTeX X documents. The plot programs support color — PROF1 uses the "pen" number as follows:

- 1 black
- 2 red
- 3 blue
- 4 green
- 5 yellow
- 6 cyan
- 7 magenta

A more detailed description follows in the next sections.

At Tübingen, the programs are found at /home/rauch/tools/pladd /home/rauch/tools/plxy /home/rauch/tools/prof1 /home/rauch/tools/rtfz

All programs are self-explaining . . .

In the following sections some general comment on the programs are summarized. For details, especially for one-purpose derivates of the programs, please contact the authors.

11.1 PLADD

The program *PLADD* may be used to interactively co-add spectra with individual weights. Parts of the spectra which are obviously corrupted, e.g. by cosmics, can be eliminated. The program expects input spectra data file with the convention ¡name¿.ii, where ii is a two digit integer (default is 01). In the case of echelle spectra, ii is the order number:

Example file names to be co-added:

• longslit

star1_date1.01

 $star1_date2.01$

star1_date3.01

star1_date4.01

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 $star1_date5.01$

• echelle

star1_date1.44

star1_date1.45

star1_date1.46

star1_date1.47

star1_date1.48

star1_date1.49

star1_date2.44

 $star1_date2.45$

star1_date2.46

star1_date2.47

 $star1_date2.48$

star1_date2.49

With *PLADD*, normalized as well as absolute flux spectra can be co-added. The absolute fluxes can be pre-normalized (with a constant value) for the processing. A Savitzky-Golay filter Savitzky & Golay (1964) may be invoked to clarify in case of heavy over-sampling.

11.2 PROF1

For the direct evaluation and visualization of line profile plot data files which are created by LINE1, or are available in WRPLOT format, the program PROF1 is available.

PROF1 can plot the data as well as process them in order to compare them to observations etc. A small "help" menu is available which can be invoked from the *PROF1* main menu. Several submenus can be accessed from this main menu (input "L" at the right row . . .). It is worthwhile to note that there is a plot box menu which can be accessed from the first row of the main menu.

For all further questions and assistance the authors are available via email.

In the following, the format of the line profile and spectra files is briefly summarized:

11.2.1 Line Profile Data Files

The line profile data files consist out of a *LINE1* header and any number of plot data sets. The header has the following structure:

A5 program code of the line profile calculation, e.g. LINE1

A8 date of the line profile calculation

A8 time the line profile calculation

A5 program code of the model atmosphere calculation, e.g. PRO2

A8 date of the model atmosphere calculation

A8 time of the model atmosphere calculation

15 number of the following plot data sets

Every plot data set is devided in three parts, i.e. a header, the plot data, and information about blends and the line broadening theories used for the calculation of the theoretical line profiles.

The header has the following format:

```
A80 comment (free)
A80 comment (free)
A80 comment (free)
A27,F11.3,A42 both strings (free) number field: central wavelength in Å
A80 title
A80 abscissa label
A80 ordinate label
I5 number of following values in the data set
```

The central wavelength in the marked field is of special importance because PROF1 uses this value as origin of an internal coordinate system. The data of the line profiles is given relative to the line center. All following program steps refer to this wavelength.

It follows the number of x-y pairs as announced in the header. There is no special format necessary but the first four columns have to be blank.

At the end of every data set an information segment follows which is used by PROF1 to find the position of blends in the calculated line profile and to get informations about the line broadening theories which were used to calculate the line profiles. This segment has the following form:

```
total number of lines (incl. blends) in this data set
A20,2F13.3 type of lines in TMAP Code (Sect. 2.1),
distance (Å) relative to the central wavelength (see above),
central depression of the line
total number of lines
A20,A10 line broadening theory used to calculate the line (e.g. VCS 88)
```

11.2.2 Spectra

The basic format of all spectra files which are accepted by *PROF1* is:

```
A20 name of spectrum

16 number of x-y pairs

6X,* x-y pairs
:
```

The number of x-y pairs is currently limited to 18000.

11.2.3 PROF1 Auxiliary Files

The program PROF1 can create a variety of auxiliary files. These files are explained in the following. They can be recognized by their extension.

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• .AEQ

This file is created if equivalent widths are measured with PROF1. It contains information about the measured interval, theoretical, and observed equivalent width (measure from the spectrum).

• .AUX

This file is created if labels are interactively inserted and saved! It can be easily edited and used for a following PROF1 session.

• .CMD

While a *PROF1* session is active, all commands are recorded and can be saved at the end of the session. Thus, a batch job is created which can be run as a procedure and creates automatically plot files. If different plot data files are copied to the same path subsequently, this procedure can create "standard" plots of different plot files.

• .DAT_REN

This file is created if a spectrum is normalized by *PROF1* and saved (basic format, see above).

• .LOAD

In this file all spectra a listed which shall be automatically loaded by PROF1 at the beginning of the session. It can be created from the "spectra menu" of PROF1.

• .MRK

In this file the base points for the rectification of spectra with natural splines can be saved.

• .REN

This file contains information about the normalization of a spectrum. Sometimes it appears necessary to correct the position of the local continuum. The following session can refer to this file, and thus, uses the same base points for normalization.

• .REN_DAT

This file contains a re-normalized spectrum.

• .SET

In the file information (defaults) about the size of the plot box etc. are saved. It is read by PROF1 at the beginning of the session.

11.3 PLXY

The program PLXY is an universal plot program for VMS and UNIX. The UNIX version is based on the PGPLOT graphics subroutine library which is provided by the California Institute of Technology (in the www, a documentation about this library is found under http://astro.caltech.edu/~tjp/pgplot/#documentation). It expects in the plot data file only a x-y table. Optional, commands can be given either by keywords (see below) in the plot data file or interactively.

11.3.1 Keywords for Commands in Plot Data Files

The program *PLXY* accepts some commands given in the plot data file which can be inserted at every position but flushed to the left. Valid are the last commands given before the end of a data set. Strings, e.g. for a comment, start directly after the ":" — without any ".".

- 0: origin of the plot $(X_0, Y_0 \text{ in cm})$, new plot box
- A: abscissa label, OFF: no label
- B: width of plot box given in cm
- C: color index, 16 colors are predefined (Fig. 1), they can be changed interactively
- D: line style
 - 1 full line
 - 2 dashed
 - 3 dot-dash-dot-dash
 - 4 dotted
 - 5 dash-dot-dot
- E: distance of tick marks
- H: height of plot box given in cm
- K: comment
- M: star position for sky map, Example:
- M: 1 0 03 44.0 -44 04 12.0 Longmore 4
- N: new data set
- 0: ordinate label, OFF: no label
- P: thickness of line
- S: plot symbols, numbers taken from the *PGPLOT* symbols, a small selection is summarized in Tab.2
- T: title

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Table 2: Selection of plot symbols available for the use in PLXY

#	symbol
840	0
841	
842	\triangle
843	\Diamond
844	*
845	+
846	×
847	*

• X: $X_{\min} X_{\max}$

• Y: $Y_{\min} Y_{\max}$

11.3.2 Output devices

Three output devices can be requested:

- 0 /dev/null, i.e. no output
- 3 PostScript file plxy_xxxx.eps, xxxx is a running counter
- 5 screen

11.4 RTFZ

RTFZ may be used to normalized spectra individually. RTFZ expects a xy table with the original spectrum. Continuum points can be set manually and a cubic spline fit will be performed for selected parts of the spectrum. In order to achieve more clarity, it is possible to smooth the spectra during the rectifying process by a Savitzky-Golay filter.

11.5 TEUV

The Tübingen EUV Absorption (TEUV) tool was created to apply interstellar absorption to theoretical stellar spectra in the wavelength range $\lambda < 911\,\text{Å}$, where bound-free absorption by interstellar elements has a string influence on observed fluxes. This has to be considered when these observations are modeled. The hot and cool components of the interstellar medium are represented in the TEUV tool.

The program is intuitive to handle and controlled via a web interface (http://astro.uni-tuebingen.de/~TEUV) where all parameters of the interstellar components can be entered and the input model-flux table can be uploaded. For further assistance, the mouse can be hovered over the blue labels.

The following fields have to be selected or filled in:

• Contact data: The user has to enter name and email address. In case any error occurs, the administrator can contact the user for assistance.

- Upload model and observation: For a fast and easy adjustment of the parameters, the resulting model will be displayed. An observation that covers the wavelength range $\lambda < 911\,\text{Å}$ is necessary for comparison. Input model and observation must contain wavelength in Å and in increasing order in the first and flux in erg/cm²/sec/A in the second column.
- Normalization: For the first overview of the observation, model, and corrected model, a normalization of the theoretical models is necessary to match the observation. Therefore, the observed flux at 928 Å or at a chosen wavelength (< 928 Å) has to be entered.
- Instrumental resolution: TEUV can also apply a correction with a Gaussian profile to the theoretical models to account for instrumental resolution. The FWHM of the Gaussian has to be given in Å.
- The two components: For both components, temperature in K, radial and turbulent velocities in km/sec as well as the column densities in cm⁻² of the single ions have to be entered.

After "Upload and Submit", the theoretical model is corrected according to the given parameters. Additionally, models with a ten times larger and smaller H I and He I column densities are created. To be able to display these models, the maximum number of output wavelength points is reduced. It may take some seconds until these corrections are applied. Then, the uploaded model, observation, and corrected model are displayed. For an easier adjustment of the parameters, the user can choose to display the models with varied H I and/or He I column densities.

If the user wants to download the resulting model, the "Download your ISM_choice" button can be used. This triggers the calculation of the corrected model without normalization, for all entered wavelength points with the instrumental resolution applied. If the TEUV tool is used to determine results that are published later, the quote to cite in the acknowledgement is given at the website. The user can store the resulting file via the "Get your results file" button.

11.6 WRPLOT

WRPLOT is a program by Wolf-Rainer Hamann et al. which can be used to manipulate and visualize data. A manual for the use of this program is available at http://astro.uni-tuebingen.de/~TMAP

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A Formula Collection

In this formula collection, those formulae are summarized which are used in the programs of the Tübingen NLTE Model-Atmosphere Package (TMAP). The abbreviations for the different transitions (CBF-...) refer to those of Sect. 1. The last numbers of the section titles are the formula numbers which have to be inserted in the atomic data file (ATOMS, Sect. 2, 2.2).

A.1 CBB Transitions

A.1.1 van-Regemorter Formula (Allowed Dipole Transitions)

$$C_{
m ij} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} \left[14.5 f_{
m ij} \left(rac{E_{
m H}}{E_0}
ight)^2
ight] u_0 e^{-u_0} \Gamma(u_0)$$

$$u_0 = h
u_{
m ij}/kT, \quad E_0 = h
u_{
m ij}, \quad \Gamma(u_0) = {
m max}[ar{g}, 0.276e^{u_0}E_1(u_0)]$$

 $\bar{g} = 0.2$ for n' \neq n else = 0.7. $E_{\rm H}$ is the ionization energy of the hydrogen ground state.

$$\pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} = 5.465 \cdot 10^{-11}$$

2 input parameters: f_{ij} , \bar{g}

A.1.2 Forbidden Transitions

$$C_{
m ij} = rac{8.631 \cdot 10^{-6}}{g_{
m i} \sqrt{T}} n_{
m e} e^{-u_0} \Omega
onumber \ \Omega = \sum_{
m i=1}^{NFIT} a_{
m i} \cdot x^{i-1}
onumber$$

NFIT+1 input parameters: NFIT, a_1, \ldots, a_{NFIT}

for the effective collision strength. Is only one parameter supplied (explicitly: 1), then $\Omega = 1$ is set.

A.1.3 Hydrogen

following Mihalas

$$C_{
m ij} = 4\pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} \left(rac{E_{
m H}}{h
u_{
m ij}}
ight)^2 u_0 f_{
m ij} \left(E_1(u_0) + 0.148 u_0 E_5(u_0)
ight) \gamma$$

 $\gamma = \beta + 2(\alpha - \beta)/\Delta n$ for $\Delta n > 1$, and $\gamma = 1$ else $\beta = 3 - 1.2/n_i$, $\alpha = 1.8 - 0.4/n_i^2$ no input parameter

А.1.4 Не п

following Mihalas & Stone

$$egin{aligned} C_{
m ij} &= \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} \left(rac{E_{
m H}}{h
u_{
m ij}}
ight)^2 u_0 f_{
m ij} e\left(e^{-u_0} \ln 2 + E_1(u_0)
ight) \gamma \ & \gamma = \min\left[n_{
m i}, 1.1
ight] \cdot \min\left[\Delta n_{
m i}, n_{
m i} - rac{n-1}{\Delta n}
ight] \end{aligned}$$

no input parameter

A.1.5 He I: Allowed Transitions from its Ground State

following Mihalas & Stone

$$C_{1
m j} = 4\pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} f_{1
m j} \left(rac{E_{
m H}}{h
u_{1
m j}}
ight)^2 u_0 E_1(u_0)$$

1 input parameter: f_{1j}

A.1.6 He I: Allowed Transitions but not from its Ground State

following Mihalas & Stone

$$C_{
m ij} = 4\pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} f_{
m ij} \left(rac{E_{
m H}}{h
u_{
m lj}}
ight)^2 u_0 \left(E_1(u_0) - rac{u_0}{u_1} e^{-0.2} (E_1(u_1)
ight)$$
 $u_1 = u_0 + 0.2$

1 input parameter: f_{ij}

A.1.7 He I: Forbidden Transitions from its Ground State

following Mihalas & Stone

$$egin{aligned} C_{
m ij} &= \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} rac{a}{n_{
m eff}}^3 u_0 rac{1}{\pi {a_0}^2} \left[a_0 E_1(u_0) + a_1 u_0 e^{-b_1 u_0} x_1 + a_2 u_0 e^{-b_2 u_0} x_2
ight] \ & x_{
m i} &= rac{b_{
m i} \left(u_0 + rac{1}{c_{
m i}}
ight) + 2}{\left(u_0 + rac{1}{c_{
m i}}
ight)^3} \ & n_{
m eff} &= Z \sqrt{rac{E_{
m H}}{h
u_{
m th}}} \quad (Z=1) \end{aligned}$$

8 input parameters: a, a_i , b_i , c_i (found in tabular form in M&S; Attention: their equation A16 is wrong!)

A.1.8 He i: Forbidden Transitions between its n = 2 Sublevels

following Mihalas & Stone

$$C_{
m ij}=\pi{a_0}^2\sqrt{rac{8k}{\pi m_{
m e}}}n_{
m e}\sqrt{T}e^{-E_0/kT}\Gamma_{
m ij}(T)$$
 $E_0=h
u_{
m ij}$

$$\log \Gamma = c_0 + c_1 \log T + c_{-2} (\log T)^{-2}$$

3 input parameters: c_0 , c_1 , c_{-2} (in tabular form by M&S)

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A.1.9 Unknown Collisional Cross-Sections

$$C_{
m ij} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} e^{-u_0} \left(1 + u_0
ight)$$

no input parameter

A.1.10 Ω -Fit of 3rd Degree

like A.1.2, but:

$$\Omega = \sum_{i=1}^{4} a_i (\log T)^{i-1}$$

4 input parameters: a_1, a_2, a_3, a_4

A.1.11 Mg II: Allowed Transitions from its Ground State

following Mihalas (1972)

$$C_{
m ij} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} 4 f_{
m ij} rac{E_{
m H}}{E_0}^2 u_0 \left(a E_1(u_0) + b e^{-u_0}
ight)$$

2 input parameters: a, b

A.1.12 van-Regemorter: Combined Levels of Complex Ions

$$C_{
m ij} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} e^{-u_0} \Gamma_{
m ij}(T)$$

$$\log \Gamma_{ij}(T) = a_0 + a_1 x + a_2 x^2 + a_3 x^3, \quad x = \log kT \quad [\text{in eV}]$$

4 input parameters: a_0 , a_1 , a_2 , a_3

A.1.21 He i: Transition between its Levels with $s \leq 4$

D. Hummer, priv. comm. no input parameter

A.1.25 Ω -Fit in Temperature, General Case

equal to **DETAIL** Formula No. 25, 8 input parameters

A.1.26 Ω -Fit in log T, General Case

equal to DETAIL Formula No. 26,

$$\Omega = \sum_{\mathrm{i=1}}^{NFIT} a_{\mathrm{i}} \left(\log T - T_{1}
ight)^{i-1}$$

NFIT+2 input parameters: T1, NFIT, $a_1, ..., a_{NFIT}$

A.2 CBF Transitions

A.2.1 Hydrogen, n = 1, ..., 10

following Mihalas

$$C_{
m ik} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} e^{-h
u_{
m th}/kT} \Gamma_{
m i}(T)$$

own fit formulae for Γ , because Mihalas is restricted in temperature, no input parameter

A.2.2 He II, $n = 1, \dots, 10$

following Mihalas, like CBF1, own fit formulae for Γ , because Mihalas is restricted in temperature, no input parameter

A.2.3 He i, $n \le 15$

$$C_{
m ik} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} \sigma_0 \left\{ u_0 E_1(u_0) - rac{0.728 {u_0}^2}{u_1} E_1(u_1) - 0.189 {u_0}^2 e^{-u_0} \left(rac{2.0 + u_2}{{u_2}^3}
ight)
ight\}$$

$$u_0 = h
u_{
m th}/kT, \; u_1 = u_0 + 0.27, \; u_2 = u_0 + 1.43$$

1 input parameter: σ_0 (in tabular form by Mihalas & Stone)

A.2.4 Seaton Formula

$$C_{\mathrm{ik}} = 1.55 \cdot 10^{13} \bar{g} \sigma_0 n_{\mathrm{e}} \frac{1}{\sqrt{T}} e^{-u_0} u_0^{-1} \quad \bar{g} = [0.1, 0.2, 0.3] \quad \mathrm{for} \ Z = [1, 2, \geq 3]$$

Z = charge of the ion $\sigma_0 =$ threshold photoionization cross-section 2 input parameters: σ_0 , \bar{g}

In case that formula 4 is requested and a cross-section of 0.0 inserted, a mean cross-section (at the threshold energy) of the Opacity Project data is calculated and used as threshold cross-section (Sect. A.4).

A.2.5 Lotz Formula

$$C_{
m ik} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} P \left(rac{E_{
m H}}{E_0}
ight)^2 u_0 \left\{E_1(u_0) - rac{lpha u_0}{u_1} E_1(u_1)
ight\}$$

3 input parameters: P, α , c $u_1 = u_0 + c$

A.2.6 Mg II $3s \rightarrow Mg III 2p^6 + \epsilon$

following Mihalas (1972)

$$C_{
m ik} = \pi {a_0}^2 \sqrt{rac{8k}{\pi m_{
m e}}} n_{
m e} \sqrt{T} \left(rac{E_{
m H}}{E_0}
ight)^2 \left\{ a u_0 E_1(u_0) + b (u_0/u_1)^2 [E_1(u_1) + e^{-u_1}]
ight\} \ u_1 = u_0 + c$$

3 input parameters: a, b, c

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A.3 RBB Transitions

A.3.1 Doppler Profiles

$$\sigma_{
m ij} = rac{\sqrt{\pi}e^2}{m_{
m e}c}rac{f_{
m ij}}{\Delta
u_{
m D}}e^{-(\Delta
u/\Delta
u_{
m D})^2}$$

$$\Delta
u_{
m D} = rac{
u_0}{c} \sqrt{rac{2kT}{m_{
m ATOM}}}$$

1 input parameter: f_{ij}

A.3.2 Voigt Profiles, only Radiative Damping

$$egin{align} \sigma_{
m ij} &= rac{\sqrt{\pi}e^2}{m_{
m e}c}rac{f_{
m ij}}{\Delta
u_{
m D}}H(a,v) \ a &= rac{\Gamma}{4\pi\Delta
u_{
m D}}, \quad v &= rac{\Delta
u}{\Delta
u_{
m D}}, \quad \Gamma &= \Gamma_{
m low} + \Gamma_{
m up} \ \end{array}$$

2 input parameters: f_{ij} , Γ

A.3.3 Voigt Profiles, Radiative and Collisional Damping (Electrons) σ_{ij} like in A.3.2, but:

$$\Gamma = \Gamma_{\rm rad} + \Gamma_{\rm St}$$

$$\Gamma_{
m St} = 6.11 \cdot 10^{-5} rac{n_{
m e}}{\sqrt{T}} \left(rac{n_{
m eff}^{
m up\,2}}{z}
ight)^2$$

(Cowley 1971) 3 input parameters: f_{ij} , Γ_{rad} , $\left(\frac{n_{eff}^{up2}}{z+1}\right)^2$

A.3.4 Voigt Profiles and "Stark Wings" (Linear Stark Effect)

$$\sigma_{
m ij} = ext{max}\left[\sigma_{
m ij}^{Formel~3}, \sigma_{
m ij}^{St}
ight]$$

$$\sigma_{
m ij}^{St} = rac{0.0368 \cdot Z f_{
m ij}}{s_{
m n} z_{
m Mikro}} U \left(rac{\Delta
u Z \cdot 1.385}{s_{
m n} z_{
m Mikro}}
ight)$$

$$s_{rmn} = \left\{ n_{\mathrm{up}} \left(n_{\mathrm{up}} - 1
ight) + n_{\mathrm{low}} \left(n_{\mathrm{low}} - 1
ight)
ight\}$$

$$z_{ ext{Mikro}} = \left[\sum_{ ext{i=1}}^{NION} Z_{ ext{i}}^{rac{3}{2}} \cdot n_{ ext{i}}
ight]^{rac{2}{3}}$$

6 input parameters: f_{ij} , Γ_{rad} , $\left(\frac{n_{eff}^{up2}}{z+1}\right)^2$, Z, n_{low} , n_{up}

A.3.5 Stark Line Broadening following Dimitrijévic

like formula 3

A.4 RBF Transitions

A.4.1 Seaton Formula

$$\sigma_{
u} = \sigma_0 \left(rac{
u_{
m th}}{
u}
ight)^s \left[lpha + (1-lpha)rac{
u_{
m th}}{
u}
ight]$$

hydrogen – like :
$$\sigma_0 = 2.815 \cdot 10^{29} \frac{z^4 g_{\rm II} (\nu_{\rm th})}{n_{\rm eff}^5 \nu_{\rm th}^3}$$

with the effective principal quantum number

$$n_{ ext{eff}} = z \sqrt{rac{R}{
u_{ ext{th}}}}$$

 g_{II} is the bound-free gaunt factor z core charge of the ion 3 input parameters: σ_0, α, s

A.4.2 Seaton Formula with Gaunt Factor

$$\sigma_{
u} = \sigma_0 \left(rac{
u_{
m th}}{
u}
ight)^s \left[lpha + (1-lpha)rac{
u_{
m th}}{
u}
ight] g_{
m II}\left(x,y,z
ight)$$

6 input parameters: $\sigma_0, \alpha, s, x, y, z$

A.4.3 Koester Formula for He i

A&A,

$$\ln\left(g\sigma_{
u}
ight) = a_0 + a_1 \ln\lambda + a_2 \ln^2\lambda, \qquad \lambda[\mathring{\mathrm{A}}]$$

3 input parameters: a_0, a_1, a_2

A.4.4 Opacity-Project Photoionization Cross-Sections, Seaton tails

See Sect. A.4.10.

A.4.5 Karzas & Latter data with Gaunt Factor

Tables taken from Karzas & Latter (1961) are used to calculate the photoionization cross-sections. 3 input parameters: z_{eff} , n (principal quantum number), l (azimuthal quantum number)

A.4.10 Opacity-Project Photoionization Cross-Sections

The Opacity Project data for a level of the ion XXXX are read from the file OP_RBF_XXXX (Sect. 8.1). The programs recognize an A10 label at the begin of the data set which represents the level name in TMAP code (Sect. 2.1). For the actual frequency grid FGRID (Sect. 3), the cross-sections are interpolated or extrapolated (including possible resonances etc.). Is formula 4 requested and a cross-section of 0.0 inserted, a mean cross-section (at the threshold energy) of the Opacity Project data is calculated and used as threshold cross-section.

no input parameter

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A.4.12 DETAIL Fit Formula

$$\ln \sigma_{
u} = \sum_{\mathrm{i=0}}^{5} a_{\mathrm{i}} \cdot \left[\ln \left(rac{
u_{\mathrm{th}}}{
u}
ight)
ight]^{i}$$

6 input parameters: a_0, \ldots, a_5

A.5 RFF Transitions

None of the RFF formulae needs an input parameter.

A.5.1 Including Contributions of LTE Levels (Unsöld)

$$\sigma_{
m kk}(
u,T) = 3.694 \cdot 10^8 e^{h
u_{
m min}/kT} rac{Z^2}{
u^3 \sqrt{T}}$$

$$u_{\min} = \min[\nu, \nu_{\text{LTE}}]$$

 $\nu_{\rm LTE}$ is the ionization threshold energy of the lowest LTE level. (cf. Unsöld 1968)

A.5.2 Including Contributions of LTE Levels with Gaunt Factors

$$\sigma_{
m kk}(
u,T) = 3.694 \cdot 10^8 \left[g_{
m ff}(
u,T) (e^{h
u_{
m min}/kT} - 1)
ight] rac{Z^2}{
u^3 \sqrt{T}}$$

A.5.3 With Gaunt Factors, no LTE Contributions

$$\sigma_{
m kk}(
u,T) = 3.694 \cdot 10^8 g_{
m ff}(
u,T) rac{Z^2}{
u^3 \sqrt{T}}$$

For the free-free Gaunt factors, the default is a calculation following Mihalas (1967, ApJ 149, 169). Since these values are calculated from a fit formula within $100 \le \lambda \le 10000 \,\text{Å}$ and an extension to longer wavelengths, data (valid from submillimetre to hard X-ray wavelengths and for temperatures from $10 - 10^9 \,\text{K}$) provided by Sutherland (1998) can be chosen by input card (Sect. 5.4).

B Fundamental Constants

In all programs of the Tübingen NLTE Model-Atmosphere Package (TMAP) the values of the fundamental constants are the same. They are summarized in Tab. 3.

Table 3: Fundamental constants used in TMAP

constant	value		dimension	name
c	2.99792458	$\times 10^{+10}$	${\rm cm~s}^{-1}$	velocity of light
e	4.80320425	$\times 10^{-10}$	esu	electron charge
$F_{ m o}$	1.25	$\times 10^{-09}$	esu	Holtsmark normal field strength
h	6.62606957	$\times 10^{-27}$	erg s	Planck constant
k	1.3806488	$\times 10^{-16}$	${ m erg}~{ m K}^{-1}$	Boltzmann constant
$m_{ m e}$	9.10938291	$\times 10^{-28}$	g	electron mass
$m_{ m p}$	1.6726217776	$\times 10^{-24}$	g	proton mass
R	109737.31568539		${ m cm}^{-1}$	Rydberg constant
π	$4 \times \arctan 1$	_		

Solar Abundances 67

C Solar Abundances

Table 4: TMAP uses the following values as solar abundances^a (Asplund et al. 2009; Maiorca et al. 2014; Grevesse et al. 2015; Scott et al. 2015b,a).

Atom	Atomic	Atomic	$\log \epsilon_{\mathrm{X}}$	n(X)/n(H)	n(X)/n(He)	mass
(X)	Number	Weight	108 cV	(by number)	(by number)	fraction
H	1	1.0079	12.00	$1.000 \times 10^{+00}$	$1.175 \times 10^{+01}$	7.375×10^{-01}
He	2	4.0026	10.93	8.511×10^{-02}	$1.000 \times 10^{+00}$	2.493×10^{-01}
Li	3	6.9410	1.05	1.122×10^{-11}	1.318×10^{-10}	5.698×10^{-11}
Be	4	9.0122	1.38	2.399×10^{-11}	2.818×10^{-10}	1.582×10^{-10}
В	5	10.8110	2.70	5.012×10^{-10}	5.888×10^{-09}	3.964×10^{-09}
C	6	12.0107	8.43	2.692×10^{-04}	3.162×10^{-03}	2.365×10^{-03}
N	7	14.0067	7.83	6.761×10^{-05}	7.943×10^{-04}	6.929×10^{-04}
O	8	15.9994	8.69	4.898×10^{-04}	5.754×10^{-03}	5.733×10^{-03}
F	9	18.9984	4.40	2.512×10^{-08}	2.951×10^{-07}	3.492×10^{-07}
Ne	10	20.1797	7.93	8.511×10^{-05}	1.000×10^{-03}	1.257×10^{-03}
Na	11	22.9898	6.21	1.622×10^{-06}	1.905×10^{-05}	2.728×10^{-05}
Mg	12	24.3050	7.59	3.890×10^{-05}	4.571×10^{-04}	6.918×10^{-04}
Al	13	26.9815	6.43	2.692×10^{-06}	3.162×10^{-05}	5.314×10^{-05}
Si	14	28.0855	7.51	3.236×10^{-05}	3.802×10^{-04}	6.650×10^{-04}
P	15	30.9738	5.41	2.570×10^{-07}	3.020×10^{-06}	5.825×10^{-06}
${f S}$	16	32.0650	7.12	1.318×10^{-05}	1.549×10^{-04}	3.093×10^{-04}
Cl	17	35.4530	5.50	3.162×10^{-07}	3.715×10^{-06}	8.203×10^{-06}
Ar	18	39.9480	6.40	2.512×10^{-06}	2.951×10^{-05}	7.342×10^{-05}
K	19	39.0983	5.04	1.096×10^{-07}	1.288×10^{-06}	3.137×10^{-06}
${ m Ca}$	20	40.0780	6.32	2.089×10^{-06}	2.455×10^{-05}	6.127×10^{-05}
Sc	21	44.9559	3.16	1.445×10^{-09}	1.698×10^{-08}	4.754×10^{-08}
${ m Ti}$	22	47.8670	4.93	8.511×10^{-08}	1.000×10^{-06}	2.981×10^{-06}
V	23	50.9415	3.89	7.762×10^{-09}	9.120×10^{-08}	2.893×10^{-07}
Cr	24	51.9961	5.62	4.169×10^{-07}	4.898×10^{-06}	1.586×10^{-05}
Mn	25	54.9380	5.42	2.630×10^{-07}	3.090×10^{-06}	1.057×10^{-05}
Fe	26	55.8450	7.47	2.951×10^{-05}	3.467×10^{-04}	1.206×10^{-03}
Co	27	58.9332	4.93	8.511×10^{-08}	1.000×10^{-06}	3.670×10^{-06}
Ni	28	58.6934	6.20	1.585×10^{-06}	1.862×10^{-05}	6.806×10^{-05}
Cu	29	63.5460	4.18	1.514×10^{-08}	1.778×10^{-07}	7.037×10^{-07}
Zn	30	65.4090	4.56	3.631×10^{-08}	4.266×10^{-07}	1.738×10^{-06}
Ga	31	69.7230	3.02	1.047×10^{-09}	1.230×10^{-08}	5.342×10^{-08}
Ge	32	72.6400	3.63	4.266×10^{-09}	5.012×10^{-08}	2.267×10^{-07}
As^a	33	74.9216	2.30	1.995×10^{-10}	2.344×10^{-09}	1.094×10^{-08}
$\mathrm{Se^{a}}$	34	78.9600	3.34	2.188×10^{-09}	2.570×10^{-08}	1.264×10^{-07}
${ m Br^a}$	35	79.9040	2.54	3.467×10^{-10}	4.074×10^{-09}	2.027×10^{-08}
Kr	36	83.7960	3.25	1.778×10^{-09}	2.089×10^{-08}	1.090×10^{-07}
Rb	37	85.4678	2.47	2.951×10^{-10}	3.467×10^{-09}	1.846×10^{-08}
Sr	38	87.6200	2.83	6.761×10^{-10}	7.943×10^{-09}	4.334×10^{-08}
Y	39	88.9059	2.21	1.622×10^{-10}	1.905×10^{-09}	1.055×10^{-08}
Zr	40	91.2240	2.59	3.890×10^{-10}	4.571×10^{-09}	2.597×10^{-08}
Nb	41	92.9064	1.47	2.951×10^{-11}	3.467×10^{-10}	2.006×10^{-09}
Mo	42	95.9400	1.88	7.586×10^{-11}	8.913×10^{-10}	5.325×10^{-09}

^a: from meteorites only

Table 4 continued.

Atom	Atomic	Atomic	$\log \epsilon_{\mathrm{X}}$	n(X)/n(H)	n(X)/n(He)	mass
(X)	Number	Weight	0 11	(by number)	(by number)	fraction
Ru	44	101.0700	1.75	5.623×10^{-11}	6.607×10^{-10}	4.159×10^{-09}
Rh	45	102.9055	0.89	7.762×10^{-12}	9.120×10^{-11}	5.845×10^{-10}
Pd	46	106.4200	1.55	3.548×10^{-11}	4.169×10^{-10}	2.763×10^{-09}
Ag	47	107.8682	0.96	9.120×10^{-12}	1.072×10^{-10}	7.198×10^{-10}
$\operatorname{Cd}^{\operatorname{a}}$	48	112.4110	1.77	5.888×10^{-11}	6.918×10^{-10}	4.843×10^{-09}
In	49	114.8180	0.80	6.310×10^{-12}	7.413×10^{-11}	5.301×10^{-10}
Sn	50	118.7100	2.02	1.047×10^{-10}	1.230×10^{-09}	9.095×10^{-09}
$\mathrm{Sb^{a}}$	51	121.7600	1.01	1.023×10^{-11}	1.202×10^{-10}	9.116×10^{-10}
$\mathrm{Te^{a}}$	52	127.6000	2.18	1.514×10^{-10}	1.778×10^{-09}	1.413×10^{-08}
I	53	126.9045	1.55	3.548×10^{-11}	4.169×10^{-10}	3.295×10^{-09}
Xe	54	131.2930	2.24	1.738×10^{-10}	2.042×10^{-09}	1.669×10^{-08}
Cs	55	132.9055	1.08	1.202×10^{-11}	1.413×10^{-10}	1.169×10^{-09}
Ba	56	137.3270	2.25	1.778×10^{-10}	2.089×10^{-09}	1.787×10^{-08}
La	57	138.9055	1.11	1.288×10^{-11}	1.514×10^{-10}	1.309×10^{-09}
Ce	58	140.1160	1.58	3.802×10^{-11}	4.467×10^{-10}	3.898×10^{-09}
Pr	59	140.9077	0.72	5.248×10^{-12}	6.166×10^{-11}	5.411×10^{-10}
Nd	60	144.2400	1.42	2.630×10^{-11}	3.090×10^{-10}	2.776×10^{-09}
Sm	62	150.3600	0.95	8.913×10^{-12}	1.047×10^{-10}	9.805×10^{-10}
Eu	63	151.9640	0.52	3.311×10^{-12}	3.890×10^{-11}	3.682×10^{-10}
Gd	64	157.2500	1.08	1.202×10^{-11}	1.413×10^{-10}	1.383×10^{-09}
Tb	65	158.9253	0.31	2.042×10^{-12}	2.399×10^{-11}	2.374×10^{-10}
Dy	66	162.5000	1.10	1.259×10^{-11}	1.479×10^{-10}	1.497×10^{-09}
Но	67	164.9303	0.48	3.020×10^{-12}	3.548×10^{-11}	3.644×10^{-10}
Er	68	167.2590	0.93	8.511×10^{-12}	1.000×10^{-10}	1.042×10^{-09}
Tm	69	168.9342	0.11	1.288×10^{-12}	1.514×10^{-11}	1.592×10^{-10}
Yb	70	173.0400	0.85	7.079×10^{-12}	8.318×10^{-11}	8.963×10^{-10}
Lu	71	174.9670	0.10	1.259×10^{-12}	1.479×10^{-11}	1.612×10^{-10}
Hf	72	178.4900	0.85	7.079×10^{-12}	8.318×10^{-11}	9.245×10^{-10}
$\mathrm{Ta^a}$	73	180.9479	-0.12	7.586×10^{-13}		1.004×10^{-10}
W	74	183.8400	0.83	6.761×10^{-12}	7.943×10^{-11}	9.094×10^{-10}
Re^{a}	75	186.2070	0.26	1.820×10^{-12}	2.138×10^{-11}	2.479×10^{-10}
Os	76	190.2300	1.40	2.512×10^{-11}	2.951×10^{-10}	3.496×10^{-09}
Ir	77	192.2170	1.42	2.630×10^{-11}	3.090×10^{-10}	3.699×10^{-09}
$\mathrm{Pt^{a}}$	78	195.0780	1.62	4.169×10^{-11}	4.898×10^{-10}	5.950×10^{-09}
Au	79	196.9666	0.91	8.128×10^{-12}	9.550×10^{-11}	1.171×10^{-09}
$\mathrm{Hg^{a}}$	80	200.5900	1.17	1.479×10^{-11}	1.738×10^{-10}	2.171×10^{-09}
Tl	81	204.3833	0.90	7.943×10^{-12}	9.333×10^{-11}	1.188×10^{-09}
Pb	82	207.2000	1.92	8.318×10^{-11}	9.772×10^{-10}	1.261×10^{-08}
Bi ^a	83	208.9804	0.65	4.467×10^{-12}	5.248×10^{-11}	6.830×10^{-10}
Th	90	232.0381	0.03	1.072×10^{-12}	1.259×10^{-11}	1.819×10^{-10}
Ua	92	238.0298	-0.54	2.884×10^{-13}	3.388×10^{-12}	5.023×10^{-11}

D How to calculate a NLTE model with TMAP

D.1 Basics

First a short explanation of the directory structure: user=\${LOGNAME}

```
/home/<user>/adaten is the directory for the atomic data checked by ATOMS2.
/home/<user>/bimod is the directory for the executable binaries of LTE2 and PRO2.
/home/<user>/fgrids is the directory for the frequency grids that were produced with SETF2.
/home/<user>/jobs is the directory for all the .job files (ATOMS2, SETF2, LTE2, and PRO2).
/home/<user>/models is the directory where the calculated models from LTE2 and PRO2 are.
```

For our script, we use the following definitions / shell variables:

```
setenv AA /home/<user>/adaten
setenv BI /home/<user>/bimod
setenv FF /home/<user>/fgrids
setenv JO /home/<user>/jobs
setenv MO /home/<user>/models
```

D.2 Creation of atomic data files with ATOMS2

- Change into the directory /home/<user>/jobs/atoms2 and copy the template job file to <name>.job. It needs a file <name> with the atomic data in /home/<user>/adaten. The name <name> can be chosen to contain all the elements used in the model (e.g. H+He)
- You need to edit both files, test_atmos_lte.job and test_atoms.job (e.g. with emacs). The first one is for LTE2, the second one for PRO2.
- After editing, execute the job files and create output files for both jobs with

```
<name>_atoms.job > <name>_atoms.out and <name>_atoms_lte.job > <name>_atoms_lte.out .
```

- Check both output files (with emacs) for error messages.
- The jobs create ASCII files, e.g. /home/<user>/adaten/<name>.atoms2.

D.3 Creation of a frequency grid with *SETF2*

- edit both files <name>_lte_setf2 and <name>_setf2 in /home/<user>/jobs/setf2. Adjust the temperature and make sure that it refers to <name>_lte.atoms2
- run both jobs and create output files with

```
<name>_setf2 > <name>_setf2.out and
<name>_lte_setf2 > <name>_lte_setf2.out .
```

• it writes the files <name>_<temperature>_lte.setf2 and <name>_<temperature>.setf2 to /home/<user>/fgrids

D.4 Adaption of the Parameter files

- extract the parameters from the <name>_atoms.out file with grep para <name>_atoms.out and from the output file <name>_setf2.out of the frequency grid
- if you want to save the parameters you can write them in a file with grep para <name>_atoms.out > object.out
- the parameter files for PRO2 can be found in /home/<user>/PARAMETER/pro2
- copy the existing parameter files to PARA_<name>.INC, PARA1_<name>.INC, PARA3_<name>.INC and adjust them by using the parameters from <name>_atoms.out

 NOTICE: the values in the parameter files should never be 0!
- compile with ssh aithp3 load pro2 <name> [x64/p64] (use x64 for 64 bit machines, p64 for penryn-processor (our quad core) 64 bit machines), the executable is copied to /home/<user>/bimod/pro2_<name>.Linux_[x64]

D.5 Calculation of a start model with LTE2

- edit <name>_lte.job with emacs. Adjust the temperature, log g, and the abundances of the elements used (as normalized number fraction) and check if the used files have the right name
- input files: <name>_lte.atoms2, <name>_temperature_lte.setf2
- start the job and write it into an output file with nice +19 <name>_lte.job > <name>_lte.out
- the lte model is written to /home/<user>/model/<name>/lte. The file is called temperature_logg_numberfraction element a_numberfraction element b, e.g. 0100000_7.00_1.000_0.000

D.6 Calculation of a NLTE model with *PRO2*

- edit the <name>_nlte.job in emacs. Adjust the temperature, log g and the abundances of the used elements and check if the used files have the right name
- \bullet input files: <name>.atoms2, <name>_temperature.setf2 and the LTE2 model as input model
- start the job with nice +19 <name>_nlte.job > <name>_nlte.out
- the output model is written to /home/<user>/model/<name>/

D.7 Strategy to calculate a NLTE model with PRO2

Some models calculate without any problem using the LTE2 start model and an atomic data files that contains all lines. This can be tested first. In case that this approach fails, try to follow these steps

Table 5: Steps to calculate a NLTE model with PRO2

Table 3. Steps to Calculate a NETE model with 11to2						
step	line	temperature	ATOMS	FGRID	— comment	
	transitions	correction	calcula			
1	no	no	LTE2	LTE2	very fast due to minimum NF , should work in all cases at least with a	
					REDUCE LOG CVEC -1	
					during the very first iterations	
2	no	yes	LTE2	LTE2	very fast	
3	no	yes	LTE2	PRO2	fast, using now the final frequency grid	
4	yes	yes	PRO2	PRO2	initial test with	
	v	·			ITMAX=O	
					NEWMAX=O	
					LP-PLOT OPTICALLY THICK/THIN, ITERATION: LAST, FILE ONLY	
					whether the atmosphere is optically thin at the outer boundary (at least two depth	
					points) and optically thick at the inner boundary (at least two depth points) - if this	
					is not fulfilled, recalculate the LTE2 models with slightly extended τ_{min} and/or τ_{max}	
					(be aware that the following $PRO2$ calculation will change the atmospheric structure	
					and the occupation numbers of the atomic levels and, thus, all line strengths - check	
					this $\tau = 1$ limit in the atmosphere by default after all $PRO2$ jobs (together with	
					temperature structure and spectral energy distribution)	
E	*****	*****	DDO0	$DD \cap \emptyset$	- ,	
5	yes	yes	PRO2	PRO2	test whether all lines can be considered in the temperature correction directly from	
					the outset	
					a) if numerical instabilities are encountered, try first	
					REDUCE LOG CVEC -1 or REDUCE LOG CVEC -2	
					b) if step a is not successful, use the STEP UP F-VALUES method, start with unprob-	
					lematic ions like H I and He II, followed by those that have low ionization fractions,	
					and finally those that are dominating in the line-forming region	
6	yes	yes	PRO2	PRO2	in the final model, neither a REDUCE LOG CVEC nor a UNSOELD-LUCY TEMPERATURE	
					$CORRECTION^a$ is allowed	
0			_			

^a The UNSOELD-LUCY TEMPERATURE CORRECTION may be used in interplay with the standard *PRO2* temperature correction because it yields the temperature of the inner atmosphere quickly and thus stabilizes the numerics. In the outer atmosphere, starting from the line-forming region, the UNSOELD-LUCY TEMPERATURE CORRECTION is not giving a reliable temperature stratification.

D.8 Example: A model with hydrogen and helium

 $T = 100000 \,\mathrm{K}$, $\log g = 7$ and number fractions H = 0.9 and He = 0.1

Basic H and He model atoms are provided by TMAD (http://astro.uni-tuebingen.de/~TMAD). These are combined then in \${AA}/H+He.

D.8.1 *ATOMS2*

- Copy the job file in /home/<user>/jobs/atoms2 to H+He.job. It needs a file H+He with the atomic data in /home/<user>/adaten. Edit both files the test_atmos.job and the test_atoms_lte.job (with emacs). Make sure you use the atomic data file H+He.
- After editing, execute the job file and create an output file for both jobs and check the output files for error messages.

Table 6: Example for the atoms2.job in the case of LTE and NLTE model atmosphere calculations

LTE model	NLTE model		
#!/bin/sh	#!/bin/sh		
set $+x$;. ${HOME}/.jobstart$	set +x;. \${HOME}/.jobstart		
#	#		
cat > options < <eos< td=""><td>cat > options <<eos< td=""></eos<></td></eos<>	cat > options < <eos< td=""></eos<>		
AUTO ION H I 16 16 16	AUTO ION H I 16 16 16		
AUTO ION HE II 14 32 -14	AUTO ION HE II 14 32 -14		
CBB-AUTO-FILL	CBB-AUTO-FILL		
CBF-AUTO-FILL	CBF-AUTO-FILL		
CBX-AUTO-FILL	CBX-AUTO-FILL		
RBB-AUTO-FILL (NONE)	.RBB-AUTO-FILL (NONE)		
RBF-AUTO-FILL (NOOP)	RBF-AUTO-FILL		
RDI-AUTO-FILL (NONE)	RDI-AUTO-FILL (NONE)		
END OPTIONS	END OPTIONS		
eos	eos		
#	#		
cp \${AA}/H+He PROATOM	cp \${AA}/H+He PROATOM		
expand options PROATOM > SOURCE	expand options PROATOM > SOURCE		
${BI}/atoms2{sys}$	\${BI}/atoms2\${sys}		
ls -1 ATOMS	ls -1 ATOMS		
#	#		
<pre>cp ATOMS \${AA}/H+He_lte.atoms2</pre>	cp ATOMS \${AA}/H+He.atoms2		
#	#		
<pre>set +x; \${HOME}/.jobend \${TMPDIR}</pre>	set +x; \${HOME}/.jobend \${TMPDIR}		

D.8.2 *SETF2*

• Edit both files H+He_lte_setf2 and H+He_setf2 in /home/<user>/jobs/setf2. Set the temperature on 100000K and make sure that it refers to H+He_lte.atoms2.

Table 7: Example for a setf2.job in the case of LTE and NLTE model atmosphere calculations

LTE model	NLTE model
#!/bin/sh	#!/bin/sh
set +x;. \${HOME}/.jobstart	set +x;. \${HOME}/.jobstart
#	#
TT=0100000	TT=0100000
#	#
type='_lte'	type=' '
#	#
cp ${AA}/H+He{type}.atoms2 ATOMS$	cp \${AA}/H+He\${type}.atoms2 ATOMS
#	#
${BI}/{setf2}{sys} << \%$	\${BI}/setf2\${sys} << %
\$TT	\$TT
PRINT CHECK	PRINT CHECK
1.30E+16	1.30E+16
-1	-1
10 10	10 10
%	%
#	#
if test -s FGRID	if test -s FGRID
then	then
<pre>cp FGRID \${FF}/H+He_\${TT}\${type}.setf2</pre>	<pre>cp FGRID \${FF}/H+He_\${TT}\${type}.setf2</pre>
else	else
echo 'no FGRID created'	echo 'no FGRID created'
fi	fi
#	#
<pre>set +x; \${HOME}/.jobend \${TMPDIR}</pre>	set +x; \${HOME}/.jobend \${TMPDIR}

• Run both jobs and create output files for them. The files H+He_100000(_lte).setf2 and are written to /home/<user>/fgrids.

D.8.3 Parameter files

- Extract the parameters from the H+He_atoms.out file with grep para H+He_atoms.out and from the output file H+He_setf2.out of the frequency grid If you want to save the parameters you can write them in a file with grep para H+He_atoms.out > object.out
- the parameter files for PRO2 can be found in /home/<user>/PARAMETER/pro2
- copy the existing parameter files to PARA_H+He.INC, PARA1_H+He.INC, PARA3_H+He.INC and adjust them by using the parameters from H+He_atoms.out and H+He_setf2.out.
 - NOTICE: the values in the parameter files should be never 0!
- compile with ssh aithp3 load pro2 H+He [x64/p64] (use x64 for 64 bit machines, p64 for penryn-processor (our quad core) 64 bit machines). The executable /home/<user>/bimod/pro2_H+He.Linux_[x64] is created

D.8.4 *LTE2*

- Edit H+He_lte.job with emacs. Set T=100000K, logg=7 and H=0.9 and He=0.1. The input files are H+He_lte.atoms2 and H+He_100000_lte.setf2.
- Start the job (an example is given below) and write its output into a file.

 nice +19 H+He_lte.job > H+He_lte.out The lte model is written to
 /home/<user>/model/H+He/lte. The file is called
 0100000_7.00_0.900_0.100.

```
\#!/bin/sh
set +x; $\{HOME\}/.jobstart
# do not edit the beginning of this file
## own job following ##
#
# common paths
GRP=/home/rauch/group
# user part
#
  directories etc.
#
code=lte2
 mod=H+He
type='_lte'
#
jobdir=${JO}/lte
flxdir=${JO}/lte
#
# model parameters
 H = 0.900
HE = 0.100
#
GGM=7.00
TTM = 0100000
#
name = \$ \{TTM\} \_\$ \{GGM\}
fn=\$\{mod\}
#
aa=${AA}/${fn}${type}.atoms2
ff = {FF}/{fn}_{s} {TTM} {type}. set f2
#
if test ! -s \$\{ff\}
then
  echo '{TTM}'
                     > SDATEN
  echo 'PRINT CHECK' >> SDATEN
  echo '1.30E+16'
                     >> SDATEN
  echo '1'
                     >> SDATEN
```

```
echo '20 10'
                      >> SDATEN
  cp ${aa} ATOMS
  \{BI\}/ setf2 \{ sys \} < SDATEN 1>\&1
  if test -f FGRID
  then
    cp FGRID ${ff}
    chmod 600 ${ff}
    echo 'new frequency grid created'
    ls - l \$ \{ ff \}
  else
    echo 'no new frequency grid created'
    break
  fi
  rm *
fi
MO=\$\{MO\}/\$\{mod\}/lte/\$\{name\}\_\$\{H\}\_\$\{HE\}
if test ! -s ${MO}
then
#
echo '2 1 H HE' > normH
echo '2 2 ${H} ${HE}' > normHE
HN='/home/rauch/bimod/normalize${sys} < normH 2>/dev/null 1>&1'
HEN='/home/rauch/bimod/normalize${sys} < normHE 2>/dev/null 1>&1'
cat > LDATEN << eos
T EFF ${TTM}
LOG G ${GGM}
DAMP=0.1
ITMAX=100
EPS = 1.0E - 6
TAU SCALE 86 90 -2
PRINT INTEGRATED EDDINGTON FLUX, ITERATION: LAST
PRINT TEMPERATURE CORRECTIONS, ITERATION: LAST, ALL DEPTHS
PRINT MODEL ATOMS (OVERVIEW)
ABUNDANCE H ${HN}
ABUNDANCE HE ${HEN}
eos
#
cp ${aa} ATOMS
cp ${ff} FGRID
{BI}/{code} sys {code} < LDATEN
if test -s MODELL
then
  cp MODELL ${MO}
  chmod 600 ${MO}
fi
fi
#
# do not edit the rest of this file
```

D.8.5 PRO2

- Edit the H+He_nlte.job in emacs. Adjust the temperature, $\log g$ and the abundances of the used elements as before and check if the used files have the right name. The Input files are H+He.atoms2, H+He_100000.setf2 and the model from LTE2 is the input model.
- Start the job (an example is given in table below) with nice +19 H+He_nlte.job > H+He_nlte.out
- The output model is written to /home/<user>/model/H+He/.

```
\#!/bin/sh
set +x; $\{HOME\}/.jobstart
# do not edit the beginning of this file
## own job
           following ##
#
# common paths
GRP=/home/rauch/group
# user part
#
# directories etc.
#
code=pro2
 mod=H+He
#
\mathbf{type} = 
jobdir=${HOME}/jobs/pro2
flxdir=${HOME}/jobs/pro2
IONFRAC=HE
  — model parameters
fn=\$\{mod\}
GGM = 7.00
TTM = 0140000
 H = 0.900
HE = 0.100
name = \$ \{TTM\} \_\$ \{GGM\}
#
#
aa = \{AA\}/\{fn\}\{type\}. atoms2
ff = {FF}/{fn}_{s} {TIM} {type}. set f2
```

```
#
if test ! -s \$\{ff\}
then
/home/rauch/prep_conts_man${sys} > noCONTS_MAN << eos
5.0 2000.1 0.1
eos
  echo '${TTM}'
                       > SDATEN
  echo 'PRINT CHECK' >> SDATEN
  echo '1.30E+16'
                     >> SDATEN
  echo '1'
                      >> SDATEN
  echo '4 10'
                      >> SDATEN
  cp ${aa} ATOMS
  {BI}/{setf2} { sys } < SDATEN 2>/dev/null 1>&1
  if test -f FGRID
  then
    cp FGRID ${ff}
    chmod 600 ${ff}
    echo 'new frequency grid created'
    ls - l \$ \{ ff \}
  else
    echo 'no new frequency grid created'
    break
  fi
  rm *
fi
#
name=${name}_${H}_${HE}
MI=\$\{HOME\}/models/\$\{mod\}/0100000\_7.00\_0.900\_0.100\_pro2
MO=\$\{HOME\}/models/\$\{mod\}/\$\{name\}\_pro2
#
if test -s \$\{MI\}
then
#
if test -s \$\{MO\}
then
  MI=\$\{MO\}
fi
#
cat > DATEN << eos
COMMENT: test 4 TMAP
.CHANGE ABUNDANCE H ${H}
                              MASS FRACTION
.CHANGE ABUNDANCE HE ${HE}
                               MASS FRACTION
.CHANGE LOGG $GGM
.CHANGE EFFECTIVE TEMPERATURE $TT
LAMBDA=4
OCCUPATION PROBABILITY FORMALISM FOR H1
OCCUPATION PROBABILITY FORMALISM FOR HE1
OCCUPATION PROBABILITY FORMALISM FOR HE2
```

```
OPACITY PROJECT RBF DATA: START AT EDGE
OPACITY PROJECT RBF DATA: MISSING HYDROGENIC
.STEP UP F-VALUES: MODEL-START: H1
                                      1.0E-03 1.5 1
.STEP UP F-VALUES: MODEL-START: HE1 1.0E-04 1.5 1
.STEP UP F-VALUES: MODEL-START: HE2 1.0E-03 1.5 1
ITMAX=20
.ERRSCH=1.0E-4
NEWMAX=2
.ERRNEW=1.E-8
RADIATIVE EQUILIBRIUM: DIFFERENTIAL/INTEGRAL FORM
. INNER BOUNDARY: LAMBDA-ITERATION
LINEARIZE HYDROSTATIC EQUATION
NO NEGATIVE POPULATION NUMBERS (LTE)
.SWITCH OFF LINES
DEPTH DEPENDENT LINE PROFILES, LINEARIZATION: FIRST
KANTOROVICH=2.
                 SWITCH LIMITS 0-->1, 1-->2, 2-->1 : 0.1 0.01 0.5
SOLVE STATISTICAL EQUATIONS ONLY RE-SOLVE PARTICLE CONSERVATION
.NO TEMPERATURE CORRECTION
. UNSOELD-LUCY TEMPERATURE CORRECTION DAMP=0.1
.UNSOELD-LUCY PARAMETERS PRINT LIMIT 0.1 TAU-WTS 0.1 1.
.REDUCE LOG CVEC -1
.PRINT OPTIONS
PRINT MODEL ATOMS (OVERVIEW)
PRINT ABUNDANCES
PRINT MAX. REL. CORRECTIONS EVERY 1 ITERATIONS
PRINT INTEGRATED SURFACE FLUX, ITERATION: EACH
PRINT CP-TIME/ITERATION, EACH
PLOT EMERGENT FLUX, ITERATION: LAST
PRINT OUTPUT MODEL, ITERATION: LAST, DEPTH INCREMENT: 1
                                                       (STRUCTURE ONLY)
PRINT CORRECTIONS OF LAST LINEARIZATION, ITERATION: LAST, DEPTH INCREMENT: 1
LP-PLOT OPTICALLY THICK/THIN, ITERATION: LAST, FILE ONLY
PLOT IONIZATION FRACTIONS \{IONFRAC\} -8.5 2.5 -10.0 0.5
MACHINE 'hostname'
eos
#
cp ${MI} MODIN; chmod 600 MODIN
cp ${aa} ATOMS
cp ${ff} FGRID
/home/rauch/data/get_OP > /dev/null
#
$\{BI\}/$\{code\}_$\{mod\}$\{type\}$\{sys\} < DATEN
if test -s MODOUT
```

```
then
  echo "new_NLTE__model_${MO}_created"
  cp \{MI\} \{MI\}_{-} 'date +\%y-\%m-\%d\}_{-}H:\%M:\%S'
  cp MODOUT ${MO
  chmod 600 ${MO
  1s -1 ${MO
  if test -s STOP
  then
    cp STOP ${MO}.converged
  fi
else
  echo "no_new_model_${MO}_created" > ${MO}.failed
  touch ${MO}.failed@$HOSTNAME
  echo "no_new_model_${MO}_created"
  if test -s MODTMP
  then
    echo "model_found_from_iteration_before_failure"
    cp ${MI} ${MI}_' date +\%y-\%m-\%d_\%H:\%M:\%S'
    cp MODTMP ${MO}_tmp
    ls - l  MO_tmp
  fi
fi
#
if test -s IONPLOT
then
  cp IONPLOT ${jobdir}/{name}. ${IONFRAC}_ion
fi
if test -s PLLP
  cp PLLP ${jobdir}/${name}.lp
fi
if test -s PLOTCORR
  cp PLOTCORR ${jobdir}/${name}.corr
if test -s PRFLUX
then
  cp PRFLUX ${flxdir}/${name}.flux
fi
if test -s STRUCTURE
then
  cp STRUCTURE ${jobdir}/${name}.T-structure
fi
#
fi
# do not edit the rest of this file
set +x; ${HOME}/.jobend ${TMPDIR}
```

D.9 Naming the *TMAP* models

In the framework of the Virtual Observatory, TMAP models and SEDs that are calculated from them have to follow a general rule. The models' names have to start with $T_{\rm eff}$ (TTTTTT) and $\log g$ (G.GG), followed by the element abundances in mass fractions. An example is

```
0100000_7.00_H:__9.0000E-01_HE:_1.0000E-01 .
```

This is suitable for a small number of elements, where the level name stays relatively short. Thus, the general form for a model name is then

```
TTTTTTT_G.GG_ABUND_<nnn>,
```

where nnn is a three digit integer code. ABUND_nnn corresponds to a file with the same name (located e.g. in the model directory). It has the form

Please use one line per element, starting with the TMAP element code. The abundances are in mass fraction. The entry for the source of the abundances is optional and format free. Avoid to use any string that will be erroneously identified as an element code by a UNIX grep command!

In the *TMAP* jobs, the following has to be inserted to set some shell variables (\${moddir} is the model directory, \${abund_old} and \${abund_new} may be used for different abundances).

```
abund_old="ABUND_001"
abund_new="ABUND_001"
#

# abundances in mass fraction (see files)
#
# old abundances
oldabund=${moddir}/${abund_old}
#
Hold='grep "H___" ${oldabund} | awk '{ print $2 }''
HEold='grep "HE__" ${oldabund} | awk '{ print $2 }''
Cold='grep "C__" ${oldabund} | awk '{ print $2 }''
Nold='grep "N__" ${oldabund} | awk '{ print $2 }''
Nold='grep "O__" ${oldabund} | awk '{ print $2 }''
NEold='grep "NE__" ${oldabund} | awk '{ print $2 }''
NEold='grep "NE_" ${oldabund} | awk '{ print $2 }''
SIold='grep "SI_" ${oldabund} | awk '{ print $2 }''
FEold='grep "FE_" ${oldabund} | awk '{ print $2 }''
FEold='grep "FE_" ${oldabund} | awk '{ print $2 }''
```

```
NIold='grep "NI_" ${oldabund} | awk '{ print $2 }''
#
# new abundances
newabund=${moddir}/${abund_new}
#
Hnew='grep "H___" ${newabund} | awk '{ print $2 }''
HEnew='grep "HE__" ${newabund} | awk '{ print $2 }''
Cnew='grep "C___" ${newabund} | awk '{ print $2 }''
Nnew='grep "N___" ${newabund} | awk '{ print $2 }''
Nnew='grep "O___" ${newabund} | awk '{ print $2 }''
NEnew='grep "NE__" ${newabund} | awk '{ print $2 }''
NEnew='grep "NE__" ${newabund} | awk '{ print $2 }''
SInew='grep "SI_" ${newabund} | awk '{ print $2 }''
FEnew='grep "FE_" ${newabund} | awk '{ print $2 }''
NInew='grep "NI_" ${newabund} | awk '{ print $2 }''
NInew='grep "NI_" ${newabund} | awk '{ print $2 }''
NInew='grep "NI_" ${newabund} | awk '{ print $2 }''
```

In the input files of e.g. PRO2 respective cards have to use the XXnew variables (XX is the element):

```
CHANGE_ABUNDANCE_H__${Hnew}_MASS FRACTION
CHANGE_ABUNDANCE_HE_${HEnew}_MASS FRACTION
CHANGE_ABUNDANCE_C_${Cnew}_MASS FRACTION
CHANGE_ABUNDANCE_N__${Nnew}_MASS FRACTION
CHANGE_ABUNDANCE_O__${Onew}_MASS FRACTION
CHANGE_ABUNDANCE_NE_${NEnew}_MASS FRACTION
CHANGE_ABUNDANCE_SI_${Sinew}_MASS FRACTION
CHANGE_ABUNDANCE_FE_${FEnew}_MASS FRACTION
CHANGE_ABUNDANCE_FE_${FEnew}_MASS FRACTION
CHANGE_ABUNDANCE_NI_${Ninew}_MASS FRACTION
```

E Colors

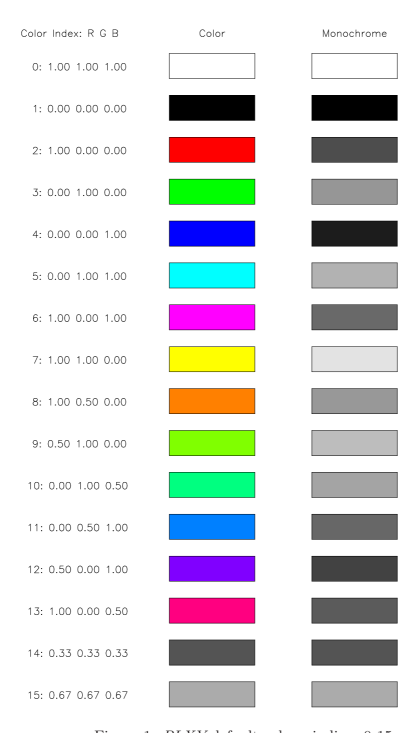


Figure 1: PLXY default colors, indices 0-15

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