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1 Introduction

The HELIX code is based on fitting the observed Stokes profiles with synthetic ones obtained from an analytic solution of the Unno-Rachkovsky equations [Unno, 1956; Rachkovsky, 1967] in a Milne-Eddington atmosphere (see e.g. Landi Degl'Innocenti [1992]). These synthetic profiles are functions of the magnetic field strength $|B|$, its inclination and azimuth, the line-of-sight velocity, the Doppler width, the damping constant, the ratio of the line center to the continuum opacity, and the slope of the source function and the source function at $\tau = 0$. The atmospheric parameters that ensure the minimum of the merit function are obtained using a very reliable genetic algorithm (PIKAIA, Charbonneau [1995]). This algorithm reaches the global minimum of the merit function with higher reliability than a classical Levenberg-Marquardt algorithm, which is also implemented in the code.

The code is able to handle an arbitrary number of atmospheric components. Atomic data files allow the flexible usage for various spectral lines.

Different versions of the code are available:

- plain IDL version: Only IDL routines are used. This version is very slow, but OS independent.
- plain FORTRAN 90 version: no IDL is required to run the code. Several architectures are supported (e.g. Linux, Solaris). This version is optimized for speed. An MPICH version is available to run the code on workstation clusters. The FORTRAN 90 version uses DISLIN to display results.
- IDL version calling FORTRAN 90 shared objects: this version allows fast inversions within the IDL environment. This is useful for analyzing the results of the inversion.

Additionally, there is an extensive IDL library available to analyze inversions of a whole map of Stokes profiles.

HELIX can be used freely. Please add a reference to Lagg *et al.* [2004] if you publish results obtained with HELIX.

2 The Physics

For a description of the physics see Lagg *et al.* [2004].

3 Obtaining / Installing the Software

The most recent version of the software is located on pulpo.mps.gwdg.de in the directory [/data/gbso/software/linefit](http://pulpo.mps.gwdg.de/data/gbso/software/linefit). This directory contains all source files, documentation and sample input files. The software is also available on the password-protected website <http://www.mps.mpg.de/homes/lagg/> under “GBSO download section”. Ask the author to get access to this web-page.

There are two methods to install the software:

- network installation: use this option if you want to use the software without changing it. This option ensures that you always use the latest release of the software. NOTE: This only works if there is a local installation (see Sect. 3.2) available on your network.

- installation on your local hard disk or your home directory: Use this option if you want to do changes in the code or if you do not have a network connection to the official distribution directory

3.1 Network Installation

The network installation will only create links to the program files located at the server at the MPS. Network-installation only works for PCs inside the MPS-network. You will find a shell-script called `net_install` in the official distribution directory. To perform the network installation please run this shell script:

```
cd /data/gbso/software/linefit ; ./net_install your_directory
```

The option `your_directory` specifies the location of the network installation. It must be a directory where you have write access. If you do not specify an installation directory your home directory will be used.

The script will install the code into the directory `your_directory/linefit`.

The script will create the necessary symbolic links to the distribution directory. It will also copy and synchronize some directories which contain user-files (such as `input wgt atom`). It creates input and output directories (`ps sav atm_archive profile_archive`).

The user has to take care that

- enough disk space is available in the output directories
- the profile directory `profile_archive` contains valid observations.

3.2 Local Installation

To install the software locally you should get the zipped tar-file `linefit_yymmdd.tar.gz` in the directory `/data/gbso/software/linefit/archive`. Here are the step-by-step instructions for the installation:

- Copy the tar-file to a location of your choice, eg. `$HOME/linefit/`:

```
cp /data/gbso/software/linefit/linefit_yymmdd.tar.gz $HOME
```
- untar / unzip the file and enter the linefit directory:

```
gunzip linefit_yymmdd.tar.gz ; tar xf linefit_yymmdd.tar
```
- create the link to the correct executable:

```
ln -sf bin/linefit.linux linefit
```

If the correct executable is not present, you have to recompile the code (see Sect. 4).
- Match the directory structure to your needs (the input file can be used to define the path to these directories):

```
$HOME/linefit/ps
```

 - directory for postscript output

```
$HOME/linefit/sav
```

 - directory for IDL-sav files

```
$HOME/linefit/profile_archive
```

 - directory or link to the spectra

```
$HOME/linefit/atm_archive
```

 - output directory for atmospheres / synthetic spectra

```
$HOME/linefit/wgt
```

 - directory for wavelength dependent weighting functions

`$HOME/linefit/atom` - directory for atomic data files

3.3 Running FORTRAN 90 version

To start the FORTRAN 90 version with the input file `default.ipt` type

```
./linefit -i default.ipt [ -x xxx -y yyy ]
```

The optional keywords `-x` and `-y` can be used to select a pixel from a map of observed profiles.

Set your environment variable `LD_LIBRARY_PATH` to include the library paths of the FORTRAN 90 compiler, the DISLIN-distribution and the shared object files (IDL-version):

```
export LD_LIBRARY_PATH=./lib:./dislin:./cfitsio:$LD_LIBRARY_PATH
```

The distribution contains the dislin-libraries for 32-bit Intel processors. If the code is run on other architectures (e.g. the sparc helios.mps.mpg.de) the library path has to be set to the according location on this machine.

If error messages appear (shared libraries missing, ...) try to recompile the code (see Sect. 4.1).

Make sure that the data referred in the input file are at the appropriate location (input-file variable `PROFILE_ARCHIVE`)

3.4 Running IDL version

The IDL-version is the original version of the software. Since it is slower and not MPI-capable the main distribution is the FORTRAN 90 version. The IDL version should not differ in results from the Fortran 90 version.

Run the startup-script:

```
./run
```

If error messages appear, you may have to adapt the startup script `run` (eg. `emacs run`). You will have to change the location of the idl software (default: `/opt/rsi/idl`).

To run the IDL-version you need to compile the FORTRAN 90 code as shared objects (see Sect. 4.2).

Note: The shared-object libraries called by IDL only work if IDL runs in 32-bit mode. You can start IDL in 32-bit mode with the command `idl -32`.

Now you should be able to use the software. Type `linefit` at your IDL-prompt and the software should start with a selection list of the 5 most recent input files.

If an error message like this appears:

```
% CALL_EXTERNAL: Error loading sharable executable.
```

you have to set the `LD_LIBRARY_PATH` environment variable before you start idl. For the korn or bash shell the command is:

```
export LD_LIBRARY_PATH='./lib:./dislin:./cfitsio:$LD_LIBRARY_PATH'
```

which sets the `LD_LIBRARY_PATH` to the `lib` directory which is in your linefit home directory and adds the initial value of `LD_LIBRARY_PATH`.

The libraries contained in the `./lib` directory are for the Intel Fortran Compiler, version 8. If you use a different compiler for creating the shared object files, you will have to change these files accordingly.

4 Compiling the Code

Note: The code is already compiled for Linux. Usually, there is no need to recompile the code. If you want to run the code on other operating systems or you want to implement new features, you will have to recompile the code.

The compilation of the FORTRAN 90 code is necessary for the IDL and the FORTRAN 90 version. Both versions use the same FORTRAN 90 routine to do the calculations. The preferred FORTRAN 90 compiler is Intel Fortran (download at www.intel.com).

4.1 FORTRAN 90 version

Goto the directory `$HOME/linefit/fortran`. Run the `./configure` script-file to create a makefile according to your needs using one or several of the following options:

- `x11` - includes DISLIN display routines
- `intel` - for compilation with the Intel Fortran compiler for Linux, version 7 or 8. Can be combined with the X11 option.
- `solaris` - for compilation with the Intel Fortran compiler 8.1 for Sun Solaris
- `solaris64` - for compilation with the Intel Fortran compiler 8.1 for Sun Solaris, 64 bit version (slightly faster than 32 bit)
- `solmpi` - for compilation with the Intel Fortran compiler 8.1 for Sun Solaris, with MPICH. To be used on the helios-cluster at the MPS.
- `scali` - creates an MPI-compatible code to be run at the Linux-Cluster in Göttingen (gwdu102). Not compatible with the X11 option.
- `mpich` - creates an MPI-compatible code for a local MPI-cluster (MPICH). Combination with X11 option is possible.

Currently only Linux versions are supported. A compilation on sun or DEC machines should be possible without major problems.

Example:

```
./configure x11 intel
```

This will create the FORTRAN 90 version with X11 support (DISLIN) for a Linux-PC using the Intel compiler.

After configure type

```
make linefit
```

to create the executable and the link to the linefit root-directory.

Note: Both the DISLIN package and the MPICH package contain a function `bzero`. This creates an error message when linking the code. To avoid this, please delete the `bzero`-function from your DISLIN-distribution, as suggested by the DISLIN author Helmut Michels. Procedure:

```
cd /usr/local/dislin/lib
cp dislin-8.1.a dislin-8.1.a.old
ar -d dislin-8.1.a _xstat.o
```

and

```
cd /usr/local/dislin/
cp libdislin.a libdislin.a.old
ar -d libdislin.a _xstat.o
```

The DISLIN directory could also be in `dislin/` of your linefit root directory.

4.1.1 CFITSIO

The FORTRAN 90 version uses the library CFITSIO to read and write FITS file. This library allows the direct usage of the TIP / TIP2 data files. The CFITSIO routines are automatically compiled with the using the makefile described above.

The CFITSIO libraries used in HELIX were downloaded from <http://heasarc.gsfc.nasa.gov/docs/software/fitsio/fitsio.html>, file: `cfitsio3004bbeta.tar.gz`.

4.2 IDL version

Goto the directory `$HOME/linefit/fortran`. Run the `./configure` script-file to create a makefile according to your needs using one or several of the following options:

- `idl.dec` - IDL version for DEC UNIX
- `idl.intel` - IDL version for Linux, Intel compiler

After configure type

```
make idl
```

to create the shared object files called by the IDL-version. The shared objects will be placed in the directory `./idl.so/`.

Currently only the 32-bit version of IDL is supported. The compiler must be set to 32-bit mode. For the x86_64 architecture it was necessary to copy the libraries

```
crtbegin.o crtbeginS.o crtend.o crtendS.o libgcc.a libgcc_eh.a
```

from the 32-bit version from directory `/usr/lib/gcc/i586-suse-linux/2.95.3` and `/usr/lib64/gcc-lib/x86_64-suse-linux/3.3.5/32/` to the directory `/usr/lib64/gcc/x86_64-suse-linux/4.1.2/32/`.

5 Using the Software

The inversion is controlled using ASCII-input files. These files are located in the subdirectory `input`. An inversion using the information from a specific input-file (eg. `default.ipt`) is invoked by the command at the IDL-prompt:

```
linefit, 'default.ipt'
```

or for the FORTRAN 90-version type at the system console:

```
./linefit -i default.ipt
```

To use the MPI-version you have to setup your MPI environment (see MPICH documentation, <http://www-unix.mcs.anl.gov/mpi/mpich/>). The MPI-version is useful for the simultaneous treatment of several pixels on different CPUs. There is no speed increase for a single pixel. To start the MPI-version and run linefit on 4 processors use the command:

```
/opt/mpich/ch-p4/bin/mpirun -np 4 ./bin/linefit.mpich -i mpitest.ipt
```

See Sect. B and Sect. D.1 for details on installation and usage of the MPICH version.

5.1 Some Examples

The easiest way to start using the code is by running and editing the sample input files (a detailed description of the parameters in the input file is given in Sect. 5.2). Several examples are available:

1. **ex_synth.ipt** IDL-call: **linefit, ipt='ex_synth.ipt'**
FORTRAN 90-call: **./linefit -i ex_synth.ipt**
 Creates a synthetic profile for the He 10830 triplet of a single atmospheric component. Adds random noise to the profiles and performs an inversion. Should result in an almost perfect to the synthetic profile.
2. **ex_synth_2c.ipt** IDL-call: **linefit, ipt='ex_synth_2c.ipt'**
FORTRAN 90-call: **./linefit -i ex_synth_2c.ipt**
 Similar to (1) but with two atmospheric components, shifted 20 km/s.
3. **ex_hinode.ipt** IDL-call: **linefit, ipt='ex_hinode.ipt', x=1, y=134**
FORTRAN 90-call: **./linefit -i ex_hinode.ipt -x 1 -y 134**
 Reads in a Hinode FITS file (**./profile_archive/hinode_data**) and performs a 2-component fit (one magnetic component and a straylight component).

5.2 The Input File

The input file consists of several parts (comments, directory structure, control parameters, data set, model atmospheres, analysis method and PIKAIA parameters). The order of the lines within the input file is not important, but should be kept like that to enhance readability. All entries beginning with a semicolon **;** are treated as comments.

Flags are set by using **'1'** (=TRUE) or **'0'** (=FALSE).

The following sections describe the keywords which are available in every part of the input file:

All lines beginning with a word which is not a keyword described in the sections below is regarded as comment. This comment is read in by the linefit program and is stored in the final result. The comments in the input file following a semicolon are not stored, they are for better readability of the input file only.

The number of comment lines is unlimited.

5.2.1 Directory Structure

Here you define the input and output paths:

PS	./ps/	path for postscript output
SAV	./sav/	path for output of IDL sav-files (these files contain input information as well as the result)
PROFILE_ARCHIVE	./profile_archive/	directory containing sav-file of observation, a directory with profiles (see Sect. 9.1 for creating the directory structure) or dat-file with single spectrum (SPINOR-format), a directory containing Hinode FITS files, a single Hinode FITS File or a mask defining Hinode FITS files (e.g. <code>**SP4*.fits</code>), or the FITS file (TIP-data: cc-file) of the observation.
ATM_ARCHIVE	./atm_archive/	directory for output atmospheres
ATM_SUFFIX	v01	add a suffix to atm-directory to identify this run
WGT	./wgt/	directory for wgt-files
ATOM	./atom/	directory for atomic data files

5.2.2 Control Parameters

These parameters control the behavior during the minimization.

DISPLAY_MAP	1	IDL version only! This flag controls if the <code>xdisplay</code> -routine should be called after a successful run of an inversion. This is useful when inverting a whole map of data.
DISPLAY_PROFILE	1	The input and fit Stokes profiles are displayed for every inverted profile (X11 and IDL version).
DISPLAY_COMP	1	If 1 then the individual atmospheric components of the fitted profiles are displayed (only active when DISPLAY_PROFILE is set to 1).
SAVE_FITPROF	1	If 1 then the fitted profile is saved as an IDL-sav file (can be used as input profile).
OUTPUT	PS	controls output media: PS for postscript, all other settings for screen (IDL-version only).
VERBOSE	1	controls verbosity: 0=none, 1=normal, 2=female

WL_NUM	256		# of WL-points (for synthesis only)
WL_BIN	1		bin size for WL-binning
WL_OFF	0		wavelength calibration: offset. if WL_OFF is not equal to zero then the WL-calibration from the input file is used - the WL-calibration contained in the data file is ignored! The WL axis is computed as follows: $WL = \text{bin\#}(\text{from } 0 \text{ to } \# \text{ of WL-points } - 1) * WL_DISP + WL_OFF$. Not used for synthesis-mode (i.e. SYNTH 1).
WL_DISP	0		wavelength calibration: dispersion per wavelength bin. See WL_OFF for details.
XPOS	000	150	two-elements vector containing xmin,xmax of the observation map to be analyzed
YPOS	000	100	two-elements vector containing ymin,ymax of the observation map to be analyzed
PROFILE_LIST	list.dat		filename of profile list. The profile list contains a two-column file with the x- and the y-values of profiles to be inverted. This list overwrites the XPOS and YPOS keywords. The profile list file must be located in the directory specified with the PROFILE_ARCHIVE keyword.
STEPX	1		step size for going from xmin to xmax. This is also the step size for the averaging.
STEPY	1		step size for going from ymin to ymax. This is also the step size for the averaging.
AVERAGE	1		if set to one then the profiles of an observation are averaged over an area of size stepx,stepy
SCANSIZE	0		step-size of multiple scans within one observation
SYNTH	0		if set to one, a synthetic profiles is minimized. The synthetic profile is calculated using the atmosphere defined in the the next section.
NOISE	0.0		Add artificial random noise of level x.
SMOOTH	0	0	smoothen profiles. First value defines smooth value, second value smooth method (0=IDL-smooth function, 1= FFT Low-Pass). Only available in IDL-version!

MEDIAN	0	apply median filter to input profiles. Helpful for removing spikes in observed profiles. Uses the IDL median function, a similar function is implemented in the FORTRAN 90 version.
MIN_QUV	0.	Define minimum magnetic signal. If the average of the 4 largest values in the magnetic signal $\sqrt{Q^2 + U^2 + V^2}$ over the wavelength range where the weighting functions are non-zero is smaller than MIN_QUV then the magnetic field is set to 0 and is not fitted.
SLIT_ORIENTATION	94.0	slit-orientation of observation (used for azimuth correction in the display routines).
HIN_SCANNR	0	Scan-number for Hinode data files (identifies the scan number for short, repetitive scans).

5.2.4 Post Processing Options

This section of the input file contains the parameters on how the profile is changed after the synthesis. A good example for such post processing parameters is the convolution with a telescope filter function. If a synthetic spectrum should be compared with an observed spectrum, it is important that the synthetic spectrum is “sent through the telescope”. If this is not done, then you compare a clean synthetic spectrum with a spectrum which is affected by, e.g., the filter curve of a Fabry-Perot.

At the moment the convolution with an arbitrary filter function is possible. This convolution is described by three parameters:

CONV_FUNC	./convolve/fp_630.dat	ASCII-file containing the filter function. Two columns in the ASCII file represent wavelength and filter transmission.
CONV_NWL	256	# of wavelength points for the filter function

The filtering is done by applying an FFT convolution. Some points are important in order to do a correct convolution with the filter curve:

- Since the filter function might have contributions at wavelengths outside the WL-range interesting for the analysis (defined with the **WL_RANGE** keyword, by the weighting function or by the observation itself), it is essential that the WL-range for the calculation of the profile is extended.
- This extension of the WL-range is done using the **WL_RANGE** keyword. Internally the code calculates a profile over this WL-range, whereas the comparison between observation and fit and the output is done using the original WL-range defined via the observation.
- The filter function is interpolated to match the spacing of the observation. It is highly recommended that you use the same WL-binning for the observation and the convolution by using the keyword **CONV_NWL**. A warning message will be issued if the WL-bins between observation and convolution function differ by more than 5 mÅ.

- For irregular gridded observation (e.g. filtergramms with 5 or 6 wavelength positions) try to find a WL-binning for the convolution which fulfills 2 criteria:
 - WL-bins of convolution function as close as possible to the observed WL-bins and
 - number of WL-bins for the convolution should be as small as possible (computing time).

To find the optimum WL-binning for the convolution you can use the IDL-tool `find_opt_binning`. This tool displays the difference between the binning of the observation and the equally spaced binning used for the convolution as a function of the number of convolution wavelength bins. A good choice for the binning is where this difference is as small as possible. The IDL command is:

```
find_opt_binning,wlin=[WL-vector of data],wlr=[6171.0,6175.5]
```

5.2.5 Model Atmospheres

Here the model atmosphere is defined. An atmosphere describes the conditions on the Sun where the line is formed. One pixel can contain more than one atmosphere: straylight from the surroundings can affect the signal at the position of the pixel and / or unresolved structures can lead to a linear superposition of profiles.

HELIX can be used in four different modes. The modes are automatically set by the physical parameters of the model atmosphere:

- Gauss-mode:
`BFIEL, AZIMU, GAMMA, VELOS, WIDTH, AMPLI, SGRAD, ALPHA`
- Voigt-mode:
`BFIEL, AZIMU, GAMMA, VELOS, VDAMP, VDOPP, SGRAD, EZERO, ALPHA`
- Voigt-Gdamp mode: similar to voigt-mode, but instead of the damping constant `DAMP` (a) the damping factor `GDAMP` (Γ) in units of the velocity of light (as defined in *Balasubramaniam and West* [1991]) is used:
`BFIEL, AZIMU, GAMMA, VELOS, GDAMP, VDOPP, SGRAD, EZERO, ALPHA`
$$a = \Gamma \frac{\lambda_0^2}{4\pi\Delta\lambda_D}$$
- Voigt-S₀ mode: same as Voigt-Gdamp mode. The only difference is that the source function at $\tau = 0$ is a free parameter. This allows fitting of profiles which are not normalized to its local continuum level. The fit parameters are:
`BFIEL, AZIMU, GAMMA, VELOS, GDAMP, VDOPP, SZERO, SGRAD, EZERO, ALPHA`
$$a = \Gamma \frac{\lambda_0^2}{4\pi\Delta\lambda_D}$$
- Voigt-physical mode, see *Balasubramaniam and West* [1991]:
`BFIEL, AZIMU, GAMMA, VELOS, GDAMP, VMICI, DENSP, TEMPE, SGRAD, ALPHA`

Every parameter of an atmosphere has the following form:

```
PAR_NAME      value min   max   fit   ;comment
```

The name of the parameter is followed by its initial value (important for synthesis), an allowed range (min to max) for this parameter and a fit-flag. The values for fit-flags are:

- 0: do not fit this parameter - use initial value

- not zero: treat this parameter as free parameter
- negative number: couple this parameter with other parameters of same negative number. This feature is useful to couple for example the magnetic field inclination angle for between two magnetic components.

Note: The coupling only works between the same parameter of different atmospheric components. **The parameters are coupled according to the ratios defined by the initial values. The ratio of the initial values is preserved by the coupling!**

NAME	Value	MIN	MAX	FIT	Comment
NCOMP	1				number of atmospheric components
BFIEL	200	0.0	2000	1	magnetic field value in Gauss
AZIMU	0.0	-90	90	-2	azimuthal angle of B-vector in degrees
GAMMA	0.0	0	180	-3	inclination angle of B-vector in degrees
VELOS	0.0	-7000	40000	1	line-of-sight velocity in m/s
WIDTH	0.20	0.10	0.70	1	line width (Gauss profile only!)
AMPLI	0.5	0.0	1.0	1	amplitude of line (Gauss profile only!)
VDAMP	0.1	0.05	10.0	1	damping constant (Voigt profile only!)
VDOPP	0.05	0.0	1.0	1	doppler broadening (Voigt profile only!)
EZERO	1.0	0.0	20.0	1	amplitude of components of propagation matrix (Voigt profile only!)
GDAMP	0.1	0.05	10.0	1	damping constant (Voigt profile, physical units mode only!), see <i>Balasubramaniam and West</i> [1991]
VMICI	0.1	0.0	1000.01		micro turbulence in m/s (Voigt profile, physical units mode only!), see <i>Balasubramaniam and West</i> [1991]
DENSP	0.1	0.0	100.0	1	density parameter (Voigt profile, physical units mode only!), see <i>Balasubramaniam and West</i> [1991]
TEMPE	10000.0.0		20000.0		temperature (Voigt profile, physical units mode only!), see <i>Balasubramaniam and West</i> [1991]
SZERO	0.02	0.00	0.04	1	source function at $\tau = 0$
SGRAD	1.0	0.9	1.1	1	gradient of source (Planck) function

ALPHA	0.5	0.0	1.0	1	Filling factor for this component (useful only with multiple components). If the filling factor is not fitted, then the value of SGRAD is used to calculate the filling factor. Note that the filling factor of all components is normalized to one before the inversion.
--------------	------------	------------	------------	----------	---

IMPORTANT: If the filling factor for one component is a free parameter, then the filling factor for at least another component must also be a free parameter. Otherwise the code has no possibility to adjust the filling factor!

USE_ATOM	he1083.0.dat	atomic data file(s) to be used for this component (more than 1 file possible). See Sect. 5.5.
USE_LINE	He	OBSOLETE KEYWORD! USE USE_ATOM INSTEAD! Lines to be used for this component.

For a two-component atmosphere you have to set **NCOMP=2** and duplicate the parameter block in the input file. It makes sense to have different ranges for the different components. You can for example use component #1 as a slow component (velocity range -7000 to +7000 m/s) and component #2 as a fast downflow component (velocity range +5000 to +40000 km/s).

5.2.6 Line Dependent Parameters

Under certain circumstances it is necessary to fit the some parameters of the atomic line. This is the case for example

- if the wavelength of the line is not known accurately enough
- if the ratio of two lines is not equal to the ratio of the corresponding loggf values (i.e. the line is formed at different temperatures)
- if a velocity gradient with height leads to different doppler shifts because of different formation heights of the lines.

Line parameters can be specified in the input file for every spectral line contained in the used atomic data files. Every line is identified by its wavelength (**LINE_ID** keyword). Available line parameters are **LINE_STRENGTH** and **LINE_WLSHIFT**. The syntax of the input file entries is similar to the atomic parameters. The first value denotes the initial value (also used for synthesis), the second and third value give the fit range, the fourth value decides whether the parameter is free or not. The same coupling mechanism as for atmospheric parameters is working (coupling using negative values for the fourth parameter).

LINE_ID	6301.5012	wavelength to identify the line
; NAME	Value MIN MAX FIT	Comment
LINE_STRENGTH	1.00 0.80 1.20 1	factor to adjust line strength
LINE_WLSHIFT	0.00 -0.20 0.20 1	adjust central WL of line (in Å)

5.2.7 Telluric Blends

Telluric blends can be fitted to the spectrum with a Voigt profile. The parameters of the blends can be specified similar to the parameters of the atmospheric components. Multi-iteration and coupling for blend parameters is disabled, since it does not really make sense here.

To limit the number of fit parameters, the blend values can be determined once (e.g. in the quiet sun or in the flat field), and then be used as predefined values (set fit value to 0), or with only very small ranges.

The telluric blend is fitted to the I profile. Since the telluric blend reduces the light level at a specific wavelength, the Q , U , and V profiles are also affected. HELIX takes this into account by also reducing Q , U , and V by the same amount as the I profile is reduced.

NBLEND	1				number of telluric blends
; NAME	Value	MIN	MAX	FIT	Comment
BLEND_WL	10832	10831	10833	1	wavelength and possible range for blend
BLEND_WIDTH	0.3	0.0	1.0	1	width of voigt-profile for blend
BLEND_DAMP	0.3	0.0	1.0	1	damping of voigt-profile for blend
BLEND_A0	0.5	0.0	1.0	1	amplitude of voigt-profile for blend (blend is not used if A0=0)

5.2.8 General Fit Parameters

This section of the input file contains general fit parameters, i.e. parameters which affect the general shape of the profile.

CCORR	1.0	0.95	1.05	1	Continuum Correction Fit. If this parameter is a fit-parameter (FIT=1), then the continuum level is adjusted within the specified range. This is useful for data where the continuum cannot be determined reliably.
STRAYLIGHT	0.0	0.0	1.0	1	Straylight correction. If this parameter is a fit-parameter (FIT=1), then a non-polarized, non-dispersive straylight is added. It will add a constant background to the I profile, and it will not affect Q , U and V .

Note that the straylight correction does not change the filling factor of the atmospheric components. However, if the code determines a straylight correction of eg. 0.2 then 20% of the light is unpolarized straylight (originating for example from instrumental effects), and only 80% of the light are of solar origin.

5.2.9 Analysis Method

Here you specify whether an approximation for the calculation of the magnetic field direction should be used or not. The approximation used is based on *Auer et al.* [1977].

APPROX_AZI	1				use approximation to calculate magnetic field azimuthal angle directly from Q and U profile [<i>Auer et al.</i> , 1977]
APPROX_DIR	1				use approximation for magnetic field direction. This keyword sets APPROX_AZI to 1. This approximation for the inclination is not very accurate for low B and high inclinations
APPROX_DEV_INC	0.				The approximation for the inclination angle of the magnetic field direction is used as an initial value for the minimization. The value is allowed to vary $\pm X^\circ$ around the approximation.
APPROX_DEV_AZI	0.				Same for azimuthal angle.
IQUV_WEIGHT	1.	1.	1.	1.	4-element vector defining relative weighting of IQUV (in that order). Additionally the code does a weighting according to the weighting file defined with the WGT_FILE keyword. This weighting scheme is used in the PIKAIA fit routine. See Sect. 5.6.
WGT_FILE	he_default.wgt				file with WL-dependent weighting function for IQUV, can be different for multiple iterations
PROFILE	voigt				functional form for π - and σ components of spectral line. Available: gauss or voigt
MAGOPT	1				include magneto-optical effects (dispersion coefficients, (Voigt profile only!))
USE_GEFF	1				use effective Landé factor (=1) or real Zeeman pattern (=0)
USE_PB	1				if set, the zeeman-splitting and strength includes the Paschen-Back effect (from the table by Socas-Navarro, see section 5.10)
PB_METHOD	1				use polynomials (=poly) or table interpolations (=table) to calculate the PB-effect

5.2.10 Pikaia Parameters

Here you control the call of the Pikaia or other minimization routines.

CODE	FORTRAN	Language to use. Available: FORTRAN (=fast) or IDL (=platform independent). The IDL code is of course only usable within the IDL interface.
METHOD	PIKAIA	Minimization method. Available: PIKAIA (=slow, but better convergence), POWELL (fast, may be trapped in local minimum), LMDIFF or PIK_LM. POWELL is available in IDL only. LMDIFF involves a Levenberg-Marquardt algorithm, based on the ODRPACK95 routines. This algorithm should be very fast and reliable. However, PIKAIA will show the best convergence. See Sect. 5.3 for details on the combined convergence strategy PIK_LM.
NCALLS	200	number of iterations in PIKAIA/POWELL/LMDIFF routine.

5.3 Convergence Strategies

PIKAIA provides a very robust technique to find the best fit to the measured Stokes vector. However, it is also very slow. The Levenberg-Marquardt based algorithm LMDIFF is approximately a factor of 40 faster than PIKAIA. Since this algorithm more likely gets trapped in local minima, the correct choice of initial parameters is important.

HELIX includes the possibility to combine the advantages of PIKAIA with the speed of LMDIFF when inverting a whole map of Stokes parameters. This is done by setting the minimization method (keyword **METHOD**) to **PIK_LM**. In a rectangular box of size **BX**×**BY** pixels the first run is done using PIKAIA for the lower left pixel of this box. All subsequent runs in this box are done using LMDIFF, with the initial parameters being the results of the PIKAIA run or any following LMDIFF run with higher fitness.

METHOD	PIK_LM	BX	BY	0.7	Use PIKAIA for the first pixel and LMDIFF for the other pixels in a box of size BX × BY . Use PIKAIA also for results with fitness lower than $0.7 \times$ the best (so far) fitness in this box.
---------------	---------------	-----------	-----------	------------	---

The fourth parameter of the keyword **METHOD** is optional and defines the fraction by how much the fitness of a LMDIFF inversion is allowed to be worse than the best fitness in the box. If the fitness is less than, e.g. 0.7, then the inversion for this pixel is repeated using PIKAIA. Setting this value to 0 (or not specifying it at all) will disable this fitness-check. Setting this value to 1 means that no LMDIFF fit is allowed to be worse than the PIKAIA fit. A higher value therefore increases computation time (since more pixels are calculated using the slow PIKAIA algorithm).

Speed Comparisons

Table 1 lists the performance of the various methods. PIKAIA clearly wins in the fitness of the derived atmospheric parameters, LMDIFF is by far the fastest method. Fig. 1 shows the resulting maps for

magnetic field strength and the velocity contour lines.

Table 1: Comparison of various convergence strategies: The box size for the combined LM_DIFF strategy was chosen to be 1×20 pixels.

method	input file entry	average fitness	time	rel. time [%]
PIKAIA	<code>METHOD PIKAIA</code>	2.26	81 h	100.0
LM_PIK	<code>METHOD PIK_LM 1 20 0.9</code>	2.17	28 h	34.6
LM_PIK	<code>METHOD PIK_LM 1 20 0.7</code>	2.18	11.2 h	13.8
LM_PIK	<code>METHOD PIK_LM 1 20 0.0</code>	2.15	6.6 h	8.1
LMDIFF	<code>METHOD LMDIFF</code>	1.82	2.9 h	3.6

5.4 Re-Write Input Files

The IDL-routine `nice_input.pro` can be used to reformat the input file. It will add the default comments and will make the file nice-looking. The calling sequence is (applied to input file `default.ipt`):

```
nice_input, 'input/default.ipt'
```

5.5 Atomic Data Files

The atomic data are defined in ascii files in the directory defined with the `ATOM` keyword. You must define at least one atomic data files for every atmospheric components by specifying the keyword `USE_ATOM` for each component. These atomic data files can be different for different components (e.g. one component uses the He-line, another component only affects the Si-line).

The format of the atomic data file is (see here the example for the He triplet at 1083 nm):

```
;atomic data file for he triplet at 1083 nm
;WL      Element Ion.   LOG_GF  ABUND  GEFF  SL  LL  JL  SU  LU  JU
10829.0911 He    1    -2.19823  0.00  2.0  1.0  0.0  1.0  1.0  1.0  0.0
10830.2501 He    1    -1.09961  0.00  1.75  1.0  0.0  1.0  1.0  1.0  1.0
10830.3397 He    1    -0.586987 0.00  1.25  1.0  0.0  1.0  1.0  1.0  2.0
```

The column entries are: wavelength, element name, ionization state (1=neutral), log-gf value (f is the oscillator strength and g is the degeneracy of the lower level), solar abundance (not used), effective Landé factor and the quantum numbers s, l and j for the lower and the upper level. The effective Landé-factor is calculated from the quantum numbers. If it differs from the value given in the atomic data file, a warning message is issued and the value from the atomic data file is used.

5.6 Weighting

The weighting function defines the wavelength dependent weighting for the I , Q , U and V Stokes vector. It is defined in a file defined with the `WGT_FILE` keyword. The file must be located in the directory defined with the `WGT` keyword (default: `./wgt/`). Multiple weighting files can be specified when running the code with multiple iterations.

The default weighting file is `he_default.wgt`:

```
;define weighting function for linefit
```

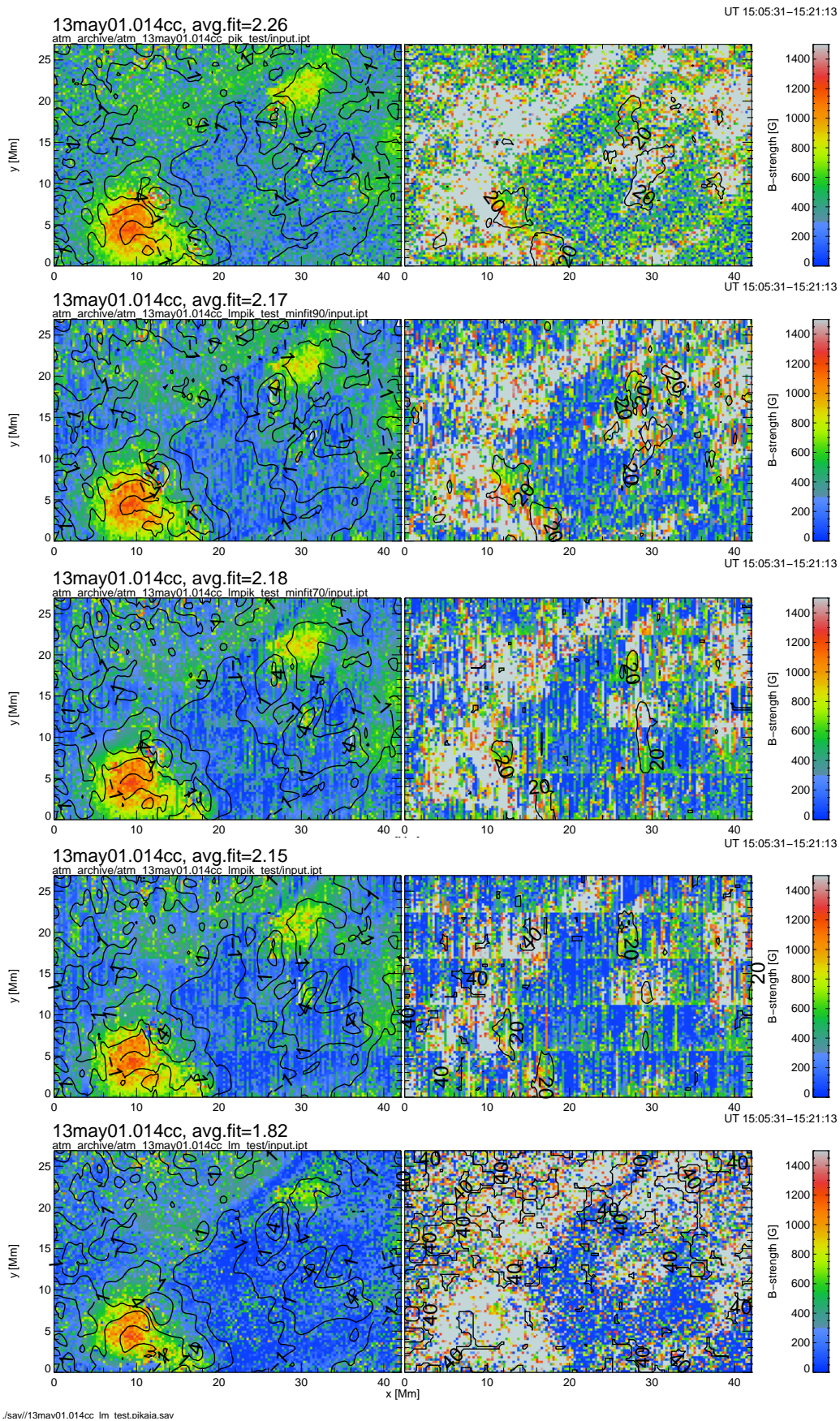


Figure 1: Magnetic field maps for the observation of an emerging flux region recorded on May 13, 2001. The results are from a two-component inversion, the methods used are in the same order as in Table 1. The contour lines show the velocity. The input file for this example can be found in Sect. E.

```

;Syntax:
;StokesPar  wgt      WLmin    WLmax
;you can define an arbitrary number of WL-ranges for every Stokes vector.
;The succeeding line will overwrite the preceding weighting function if the
;WL-ranges overlap.
I           1.0      10829.6   10831.5   ;good region for I, no telluric blend
I           0.5      10832.5   10835.0   ;include region right of telluric blend

V           0.2      10828.5   10830.0   ;V-weighting for region with Ca-Blend
V           1.0      10829.6   10832.0   ;V-weighting right of Ca-Blend

Q           0.2      10828.5   10830.0   ;Q-weighting for region with Ca-Blend
Q           1.0      10829.6   10832.0   ;Q-weighting right of Ca-Blend

U           0.2      10828.5   10830.0   ;U-weighting for region with Ca-Blend
U           1.0      10829.6   10832.0   ;U-weighting right of Ca-Blend

```

The keyword **IQUV_WEIGHT** multiplies the weighting function of every Stokes parameter with the specified value.

5.7 Correction for Scattering-Polarization

The analysis of the data set 13may01.014 showed, that the Q and U profiles can be affected by scattering-polarization. The unpolarized light beam from the solar surface hits the atoms in higher layers. The scattered light will show a linear polarization signal due to two different effects (see *Trujillo Bueno et al. [2002]*):

- the Hanle effect modifies the atomic polarization of degenerate atomic levels due to the magnetic field. The magnetic field vector has to be inclined significantly for this effect to operate
- anisotropic illumination of the atomic system (close to or above limb) will cause an imbalance in the population of the sublevel of degenerate atomic levels.

Both effects produce a linear polarization signal. The resulting Q and U signals can be used to calculate the magnetic field orientation and the magnetic field strength by applying complicated polarization diagrams (eg. *Nagendra et al. [1998]*; *Faurobert-Scholl [1992]*; *Bommier et al. [1991]*).

In the special case of a LOS-direction perpendicular to the solar surface (disk center) and a magnetic field direction parallel to the solar surface (horizontal field) the polarization signal due to the Hanle effect is oriented along the magnetic field direction:

$$\tan(2\phi) = (U/Q) \quad (1)$$

This relation gives reasonable results for horizontal fields down to a μ -angle of >0.6 (Javier Trujillo Bueno, personal communication).

To correct for this problem, we made the following approach:

- The unpolarized radiation is assumed to show a Gaussian line profile with a line width and a velocity shift calculated from the I profile.
- a Gaussian profile with a free amplitude A_0 but fixed values for width and line shift (from I) is fit to the Q and U signals simultaneously with the fit of all other free parameters of the inversion.
- the Gauss profile is distributed to Q and U according to the preferred direction (either B-field or limb direction)

- every line of a multiplet can have a separate amplitude.

To apply this correction in the inversion the following line in the 'DATA SET' part of the input file is needed:

STRAYPOL_CORR **50** **B** iteration steps and orientation of scattering-polarization correction. Orientation: 'B' = along B-field, 'X' = a number defining an angle manually

The first number gives the number of iterations for the Gauss fit to the I profile (determination of width & velocity shift). The second entry can be either 'B' for a correction according to Hanle, or a number in degrees giving the angle ϕ for a manual correction.

STRAYPOL_AMP **0.0015** This defines the straypol-amplitude to be used for a synthesis. Not used for fitting!

5.8 Multi-Iteration Method

The multi-iteration method is a technique to change the parameters relevant for the minimization after a certain number of iteration steps. The idea is to start with a simple atmospheric model to fix for example line-of-sight velocity and line broadening. In a next step a detailed analysis of the magnetic field direction can be performed.

Example: The minimization should start with the approximation for directions. After 200 iterations we consider all parameters to be determined except for the magnetic field, which should now be treated as a free parameter (no approximation anymore!). The implementation in the input file is:

NCALLS **200** **50** number of iterations in PIKAIA routine.

The first Pikaia run will have 200 iterations, the second run has 50 iterations.

Now we must change the lines defining the atmosphere:

BFIEL **200** **0.0** **2000** **1** **20** **1** magnetic field strength
AZIMU **0.0** **-90** **90** **-2** **100** **-2** azimuthal angle
GAMMA **0.0** **0** **180** **-3** **100** **-3** inclination angle

These lines do the following: The first 200 iterations will be calculated in the usual manner. The result of this run is then used as the initial guess for the second run. The second run now only changes the parameters **BFIEL**, **AZIMU** and **GAMMA**. The value for **BFIEL** is allowed to vary 20% of the initial scaling range around the new initial value. Azimuthal and inclination angle are completely free (100% variation allowed). If a second magnetic component is present, then the '-' sign would indicate a coupling between the magnetic field direction of both components.

All other atmospheric parameters are left unchanged during the last 50 iteration steps.

In order to use the approximation for the first 200 iterations but not for the last 50, we have to set the line:

APPROX_DIR **1** **0** use approx. for first run only!

The following keywords can be set differently for multiple iteration runs: **BFIEL**, **AZIMU**, **GAMMA**, **VELOS**, **WIDTH**, **AMPLI**, **VDAMP**, **VDOPP**, **ZERO**, **SGRAD**, **EZERO**, **ALPHA**, **APPROX_AZI**, **APPROX_DIR**, **APPROX_DEV_INC**, **APPROX_DEV_AZI**, **IQUV_WEIGHT** (for example, use 8 values for 2 multi-iterations), **MAGOPT**, **NCALLS**, **METHOD**, **WGT_FILE**.

5.9 Applying Different Atmospheres

With a small driver program it is possible to apply different model atmospheres for one data set depending on the results of a first run. This allows to invert the data set with a simple model and apply a more complex model only to the some pixels of the data set. This driver program is written in IDL, it therefore works only with the IDL version.

The following IDL-code is an example for such a driver program (see **driver.pro** in **idlpro/** directory):

```

;program to perform more complex linefits depending on result of
;simple linefit
pro driver

                                ;do simple linefit
linefit,ipt='emf_1comp_abs.ipt',savall=savall
restore,savall

                                ;check for parameters which make a
                                ;more detailed analysis necessary
idx=where(pikaia_result.fit.atm.par.vlos gt 5000.)

if idx(0) eq -1 then return

                                ;call more complex linefit
list=transpose([[pikaia_result.fit(idx).x],[pikaia_result.fit(idx).y]])
atm_ini=pikaia_result.fit(idx).atm
linefit,ipt='emf_2comp_abs.ipt',list=list,atm_ini=atm_ini
end

```

First, an inversion according to an input-file **emf_1comp_abs.ipt** is performed. The results of this inversion are stored in a sav-file, the name of this sav-file is returned in the variable **savall**. This sav-file is restored.

Then we check for the parameter which decides whether we want a more complex inversion or not. In this case, all points with line-of-sight velocities greater than 5000 m/s should be picked out.

A list of the x and y-pixels for these points is created (**list**). The result of the simple 1-component inversion is used as an initial guess for the more complex, 2-component run defined in the input-file **emf_2comp_abs.ipt**.

5.10 Paschen-Back Effect

The parameter **USE_PB** forces linefit to include the tabulated version of the Paschen-Back effect for the He 10830 line [Socas-Navarro *et al.*, 2004]. Two methods are implemented:

- **PB_METHOD poly** uses the polynomials to fit the deviation between the Zeeman levels with and without Paschen-Back effect [Socas-Navarro *et al.*, 2005]

- **PB_METHOD table** uses a tabulated version and a quadratic interpolation to calculate splitting and strength of Zeeman sublevels [Socas-Navarro, 2005].

5.11 Running Linefit in Batch Mode

This chapter applies to the IDL version only. The FORTRAN 90 version can easily be run in batch mode using the standard UNIX **at** command.

A small shell-script allows to run linefit within a batch-file. The shell script is in the root-directory of the linefit program and is called **batch**.

You can run several input files subsequently (nice for long weekends). The file is based on the **run** script file. The command to call several input files is:

```
./batch ipt-file1.ipt ; ./batch ipt-file2.ipt ; ./batch ipt-file3.ipt
```

It is a good idea to set the parameters **DISPLAY_PROFILE** and **DISPLAY_MAP** to 0. X-display export might not work in batch mode. Also it is recommended to set the verbosity to 0, otherwise a huge log-file might be created.

5.12 Azimuth Correction

The azimuthal angle returned by the linefit program lies between -90° and $+90^\circ$. It is not corrected for the 180° ambiguity of the Zeeman-analysis. An azimuthal angle of 0° points to the $+Q$ direction. The $+Q$ direction for TIP is always oriented along terrestrial north-south direction.

To convert this angle to the x- and y-axis of the observation the procedure **azi_qu2xy.pro** can be used. The conversion in this routine is:

- rotate the angle from the $+Q$ direction to the $+y$ -axis: $\phi_1 = \phi_Q + (\alpha_{slit} - 180)$ where α_{slit} is the slit orientation as given in the header of the observation files.
- rotate the angle to the $+x$ -axis: $\phi_{xy} = \phi_1 - 90$.

In this system an angle of 0° points to the $+x$ -axis, and $+90^\circ$ point to the $+y$ -axis.

5.13 Display results

5.13.1 Structure of output data

The results of an inversion are stored in the directory defined with the **ATM_ARCHIVE** variable (default: **./atm_archive/**). Profiles and atmospheres are stored for every single pixel of the inverted map. The filenames for the final atmospheres / fitted profiles are:

```
./atm_archive/observation_ATM_SUFFIX/x000/x000y000_atm.dat
./atm_archive/observation_ATM_SUFFIX/x000/x000y000_profile.dat
```

for pixel $x=0$ and $y=0$ of your map.

5.13.2 Prepare data for IDL display routine using `make_sav.pro`

The IDL routine `make_sav` is used to convert the directory structure returned by the FORTRAN 90 or IDL inversion (directory `./atm_archive/`) to an IDL sav-file. The usage of this routine is:

```
make_sav,data_dir='atm_archive/atm_13may01.014'
```

This will create a sav-file from the specified directory tree.

You can combine the results of different runs into one sav-file. This is recommended if you run for example a 1-component inversion for the whole data set and a 2-component inversion for a small part of this data set. The command to combine the results of two inversions is:

```
make_sav,data_dir='atm_archive/' + \
  ['atm_13may01.014_comp1','atm_13may01.014_comp2']
```

This will first read the directory `atm_archive/atm_13may01.014_comp1`. Every pixel not containing a valid fit (i.e. with `fitness=0`) will then be replaced by the results stored in the directory `atm_archive/atm_13may01.014_comp2`. The number of components in the resulting sav-file is equal to the maximum number of components of the individual inversions.

Be aware that the order of the directories is important: The pixels from the first directory are not overwritten by the results contained in the second or subsequent directory. There are 4 different methods to combine the results from different directories:

[f] best fitness: store atmospheres with best fitness.

[r] fitness ratio: overwrite atmosphere of first directory only when fitness of 2nd (3rd...) directory is better by a specified factor.

[o] overwrite: overwrite atmospheres from first directory with results from 2nd (3rd, ...) directory.

[z] zero fitness: only overwrite the atmospheres where the fitness is zero (i.e. no atmosphere present).

You will be prompted to enter the corresponding method interactively when running `make_sav`.

The filename of the newly created sav-file will be constructed from the name of the observation and the atm-suffixes of the output directories for the atmospheres (`ATM_SUFFIX` keyword).

5.13.3 Plot Fitted Profile of a Map

To plot the fitted profile from a single pixel of an inverted map you can use the IDL-routine `get_stokes.pro`. The syntax of the routine is:

```
fitprof=get_stokes(result=pikaia_result,x=0,y=0,/show,verbose=0)
```

The structure `pikaia_result` is the output of the routine `make_sav` (see Sect. 5.13.2). The keywords `x=0`, `y=0` specify the pixel of the map and the keywords `/show`, `verbose=0` control if the profile is printed to the screen and the verbosity.

The atmosphere and the input file parameters controlling the calculation of the synthetic profile are taken from the result stored in the IDL-structure `pikaia_result`.

5.13.4 Print Profile

Every plot of profiles on the screen can easily be printed by just typing **p** on the IDL command line. **p** is a shortcut to the widget **profile_settings** and opens a widget where you can change some parameters concerning the layout of the plot.

5.13.5 Widget Application **xdisplay.pro**

Once you have created the IDL sav-file you can use the display routine by calling the IDL-command:

```
xdisplay, './sav/l3may01.014.pikaia.sav'
```

xdisplay can also display the observed profiles. The observed profiles have to be in the same place as they were when the inversion was done. Additionally **xdisplay** searches for the observations in the directory **./profile_archive**. Copying the observation data to this directory (or creating a link named **profile_archive** to the observations) allows to access the observation data in other than the original directories.

The **xdisplay** program is a widget application and is more or less self explaining. Some important features are:

- Click on plot action: clicking on a pixel of the map can
 - create a postscript file of the profiles
 - perform a fit using original input-file data
 - do a fit using the file **xdefaults.ipt** (use 'edit input' button)
 - print the atmospheric parameters for a pixel to the IDL terminal
 - zoom into the window using the left mouse button
 - create line plots of parameters along a cut in the map
- select parameter maps
- sort atmospheric components
- loop tracing module

You can define several actions when clicking on the parameter map with the right mouse button:

- The plot action 'PS-Plot' will create a postscript file of the profile of the pixel you select in the map. Use the button 'Profile Settings' to define the layout of the profile.
- If the plot actions 'Fit (original)' or 'Fit (xdisplay.ipt)' are used a click on the map will call the linefit-program and redo the fit for one pixel. If the number of components of the new fit is less than the number of components of the displayed maps then you have the possibility to replace the result in the original sav-file, when the box 'Replace original data with result of new fit' is marked.
- The plot action 'Print ATM' will print out the atmospheric parameters to the IDL terminal window.

- The 'Zoom in' option allows you to select a region from the map. Hold the left mouse button down to define one edge of the region and release the mouse button at the other edge of the region. You can manually select the region by typing in the pixel numbers for the lower left (LLx and LLY) and the upper right (URx and URY) corner. 'Reset to full size' will return to the whole map.
- 'Cut' allows you to produce line plots along a cut in the parameter map. Use the mouse in the same way as for 'Zoom in' to define the start and end point of the cut. The cut will be shown for the parameters and the components displayed in the parameter maps. The cut will be displayed in the parameter maps. To remove the cut from the maps, use the 'Delete Cut' button.
- 'Save obs+fit' will create an IDL save-file from the pixel of the map you click. This save-file contains the variables `observation`, `atm`, `fit`, `wgt`, `input`, `xval`, `yval`, `icont` from this pixel. This save-file is useful for writing your own plotting routines (see Sect. 5.13.7).

The button 'All Comps.' is used to plot all available components. To select, for example, component number one set this button to 'Comp 1'. 'Min Comp' selects the component with the smaller values in every parameter map. 'Operation' allows you to perform a mathematical operation between the components. To define an operation use the file `define_operations.pro`.

The button 'Image Settings' can be used to plot data only if they fulfill special criteria ((e.g. filling factor greater than 0.3).

The button 'Display Settings' can be used to define individual ranges for the z-axis (color bars) of the different maps. A z-range of 0 to 0 indicates automatic scaling. Plot titles and character size are also defined here. A default title and subtitle are generated when leaving the title and subtitle field empty. No title will be created when a space is used as a title string. You can overplot the azimuthal angle as arrows (without arrowheads - 180° ambiguity!) by activating the 'Overplot Azimuth' button.

The button 'Profile Settings' defines the layout of the individual profiles plotted when walking with the mouse over the maps or when printing a profile using the 'PS-Plot' option of the 'Click on plot action'.

Automatic z-scaling is controlled using two options: 'Min-Max' scales from the minimum to the maximum value of the map, 'Fit Range' scales from the minimum to the maximum value defined in the input file.

The maps can be smoothed using the IDL median filter with the 'Median Maps' slider. Invalid pixels (either not calculated, calculated with error or not drawn due to 'image Settings') are usually drawn as missing data (white pixels). The 'Interpolate' button can be used to interpolate over these invalid pixels.

The 'Param sort' button controls the order of the components. If 'none' is selected the components are ordered as they come out as a result from the inversion. This button can be used to sort the components for their values. For example, selecting 'VLOS' will define the component with the highest velocity as component one.

The loop tracing module can be used to trace loops according to the magnetic field direction information. Details on the loop tracing are described in the emerging flux nature paper.

For the loop-tracing module the azimuth-correction according to section 5.12 is applied. This correction is not applied to the 2-dimensional color-maps!

5.13.6 Azimuth Ambiguity

Zeeman polarimetry has an intrinsic 180° ambiguity in the determination of the azimuth angle with respect to the line of sight direction. When converting the magnetic field vector to the solar reference frame this ambiguity problem must be solved. `xdisplay` offers the possibility to calculate the magnetic field in solar coordinates. It offers several methods to handle the 180° ambiguity problem:

- Choose the 0° solution
- Choose the 180° solution
- Choose the solution where the inclination angle to the solar normal is smallest
- Choose the solution where the inclination angle to the solar normal is largest
- Find a smooth ambiguity solution (Ambi-Smooth button).

When selecting the Ambi-Smooth button you are asked to define pixels in the map where you know the solution of the ambiguity problem (e.g. fields in the umbra should be vertical to the solar surface). You can define an arbitrary number of pixels - the more pixels you pre-define the solution the easier it gets for the code to find the correct solution. Starting from these predefined solutions the routine `los2solar.pro` searches for a smooth solution in the solar B_x , B_y , B_z maps.

Note that this solution is only a hint on how the real solution could be! More sophisticated routines (e.g. involving magnetic field extrapolations) should be used for publications.

5.13.7 Plot multiple profiles from different inversions

This example shows how to plot multiple profiles in one plot. This method is based on the save-files created with the `xdisplay` function 'Save obs+fit' (Sect. 5.13.5).

```
pro plot_result
  restore,'sav/pix_13may01.018_free_2x2_v01_x002y042.sav',/v
  obs1=observation
  fit1=fit
  wgt1=wgt

  restore,'sav/pix_13may01.018_free_2x2_v01_x014y042.sav'
  obs2=observation
  fit2=fit

  plot_profiles,obs1,obs2,fit1,fit2,title='Multiple Profiles',/init, $
    lthick=[1,2,3,4],color=[3,1,2,4],lstyle=[0,2,3,4], $
    fraction=[1,.4,.4,.4],iquv='IQUV',weight=wgt1
end
```

5.14 De-speckling of Maps

When inverting a whole map of an observation it is unavoidable that some pixels are not converting to the optimal parameters. The routine `despeckle.pro` can be used to run the linefit-procedure on these speckles. The call of the despeckle routine is:

```
despeckle, sav='savname.sav', par='inc', dev_perc=10.
```

This example runs the speckle-filter for the results of a map (stored in the sav-file `savname.sav`). The area used for despeckling is defined by clicking on the map. It checks for speckles in the parameter INC (mag. field inclination) and allows a deviation of the surrounding pixels of 10%. If the deviation is larger, then the pixel is recalculated using the original input-file parameters. If the recalculation gives a better fitness, the original result will be replaced.

The parameter `par='inc'` can be replaced by a vector of parameters in order to take into account the smoothness in more than one parameter, eg: `par=['inc','fitness','azi']`.

The despeckle routine can also be used on pixels with a low fitness. The call

```
despeckle, sav='savname.sav', minfit=1.5, input='xdisplay.ipt'
```

will re-run all pixels with a fitness of less than 1.5 using the input file `xdisplay.ipt`.

Further parameters / keywords to despeckle are:

<code>/full</code>	Apply despeckling to whole map (no selection using mouse).
<code>position=[bx,by,ux,uy]</code>	Apply despeckling to selected region.
<code>/all</code>	Apply despeckling also to pixels surrounding the 'bad' pixel.
<code>maxval=0.55</code>	If value of the parameter is larger than <code>maxval</code> the pixel is recalculated. Must have same number of elements as <code>par</code> .
<code>minval=0.20</code>	Similar to <code>maxval</code> .
<code>comp=1</code>	Analyze only the map of the atmospheric component 1. Default is to use the first component, which is component 0.

The despeckle routine displays the pixels to be recalculated on a map. If you are happy with the selection of the pixels the code writes out a list of bad pixels to `despeckle.list` which can be used in an input file with the `PROFILE_LIST` keyword. If you answer the question 'Starting linefit using ipt=...ipt for speckles [Y/N]?' with 'Y' linefit will run in IDL mode and calculate marked pixels.

6 The TIP data format

6.1 The TIP FITS file

HELIX was developed for the polarimetric data from the TIP instrument. In order to use HELIX for other data sets it is recommended to use the same FITS format. This section describes the TIP FITS format.

Important: The filename must end with the letters 'cc', indicating that the data file is calibrated and cross-talk removed. Example: `myfitsfile.fitcc`.

The header of the FITS file is as follows:

```
SIMPLE = T / FITS standard
BITPIX = 32 / 4 byte twos-compl. signed integers
NAXIS = 3 / number of axes
NAXIS1 = 1009 / lambda
NAXIS2 = 454 / y
NAXIS3 = 12 / number of pixels in x-direction
```

The example above contains the Stokes parameters in the following format:


```

NAXIS2 =                454 / y-size of cont-image
WL_NUM  =                1009 / WL-bins
WL_OFF  = 15643.52386351 / WL-Offset
WL_DISP =                0.02317366 / WL-Dispersion

```

It contains one image of size **NAXIS1**×**NAXIS2** containing the continuum level for every single profile. The header values **WL_NUM**, **WL_OFF** and **WL_DISP** contain the number of wavelength points, the starting value for the wavelength vector and the dispersion (increment for every wavelength pixel) respectively.

For an irregular wavelength grid the wavelength calibration can be defined in the first extension of the FITS file. This can be done by using the **writefits.pro** file from Solar Software:

```

IDL> wlvec=[15650.0d, 15650.2d, 15650.5d ,15650.6d, 15650.8d]
IDL> writefits,'name_of_file.001ccx',wlvec,/append

```

This will create an extension to the FITS file which will be used as the wavelength calibration. In this example the wavelength vector has 5 elements.

HELIX will always try to use the **first extension** in the FITS file as the wavelength calibration. If this information is not contained in the **.ccx** FITS file then the FITS header values **WL_OFF** and **WL_DISP** are used. **Note:** The values **WL_OFF** and **WL_DISP** in the input file have highest priority. These values will be used for the wavelength calibration if present in the input file!

7 Using Hinode Data

The example **ex_hinode.ipt** (Sect. 5.1) shows how to work with Hinode data. The code is able to directly read in the calibrated Hinode FITS files. The code needs additional information in the format of the **ccx** files described above. These files are created using **make_ccx.pro** contained in the **./idlpro/** directory of your distribution.

You can test the usage of the **make_ccx.pro** routine by applying it to the Hinode sample data set contained in the distribution:

```

IDL> make_ccx,hinode_dir='./profile_archive/hinode_data/',level_quiet=0.001

```

make_ccx performs a Voigt function fit to the iron lines to determine the wavelength calibration. For this calibration it uses an average over the 'most quiet' profiles along one slit position (all profiles with magnetic signals smaller than the value defined with the **level_quiet** keyword). It also determines the local continuum value, the average continuum value along the slit and the average continuum value for the whole image.

8 Tips & Tricks

8.1 Writing out synthetic profiles

You can write out a synthetic profile by setting the **SYNTH** keyword to 1, the **SAVE_FITPROF** to 1 and the number of iterations **NCALLS** to 0. The synthetic profile is written to **./profile_archive/ipt-name.profile.sav** (IDL-version) and **./atm_archive/ipt-name.profile.dat** (FORTRAN 90-version).

8.2 Convergence

Several fitting parameters influence the shape of the spectrum in a similar way: the damping **VDAMP**, the amplitudes of components of propagation matrix **EZERO** and the source function gradient **SGRAD** are all changing the depth of the absorption signature. Test runs showed that the convergence significantly improves when fitting only one of these parameters and setting the other two parameters to a fixed, intermediate value. Good results were achieved with the following settings:

```

VDAMP      0.35      0.00      0.70  0 ;damping constant (Voigt only)
VDOPP      0.30      0.01      0.70  1 ;doppler broadening (Voigt only)
EZERO      2.00      0.00     10.00  0 ;amplitude of components of
          ; propagation matrix (Voigt only!)
SGRAD      1.00      0.00      8.00  1 ;gradient of source function
ALPHA      1.00      0.00      0.00  0 ;Filling factor for this component

```

In this example the depth of the absorption is controlled by **SGRAD** and the width of the line by **VDOPP**. **EZERO** and **VDAMP** are fixed to 2.0 and 0.3 respectively. Remember that this setting is a trick to increase the stability of the inversion. Parameters like magnetic field or velocity are retrieved more reliably. The physical meaning of **VDAMP**, **EZERO**, **SGRAD** and **VDOPP** is changed!

8.3 Filling Factor

In a 2-component model the filling factor **ALPHA** is changing the amplitude of the 2 components relative to each other. This is very similar (but mathematically not identical!) to a change in the source function gradient **SGRAD**.

Stability of the fit greatly increases when coupling **SGRAD** for both parameters and fitting only the filling factor **ALPHA**. This is the preferred technique when fitting multiple components, especially in the He I 1083 nm line.

Another method is to set the filling factor to a fixed value and allow **SGRAD** to be free for all components. Then the ratio of **SGRAD** for the two components can be taken as a proxy for the filling factor. This behavior is already implemented in the display routine **xdisplay**. When the filling factor **ALPHA** is constant for both components, the code uses the ratio of **SGRAD** for the different components as the filling factor.

Defining both, **ALPHA** and **SGRAD** as free parameters will likely cause a non-stable determination of both parameters.

9 Helper Routines

9.1 Prepare directory structure for spectra

The IDL version of the program can directly use the sav-file created by the **line_id** data reduction routine. This is the fastest choice when working with IDL.

FORTTRAN 90 is not able to read this sav-file. In order to apply the FORTRAN 90 code to a whole TIP observations the data have to be prepared to a special subdirectory structure. This directory structure can be created automatically with the shell-script **tar_prepare**. This shell script has to be applied to a tar-file, created with the IDL tool **line_id.pro**, which is a part of the data reduction software for the TIP data. The usage is:

```
./tar_prepare /disk1/irpolar/profile_archive/13may01.014.tar
```

This command will create a target directory

`/disk1/irpolar/profile_archive/13may01.014` with the subdirectories `x000` to `x146` corresponding to the number of scanning steps of this observations.

To apply the program to this observation, the following parameters have to be set in the input-file:

```
PROFILE_ARCHIVE      /disk1/irpolar/profile_archive/  
OBSERVATION         13may01.014
```

9.2 Prepare directory structure for spectra from IDL sav-file

If you have the profiles in an IDL sav-file and this sav-file has the accepted format (Solar-MHD or linefit) then you can use the IDL routine `sav2data.pro` to create a directory structure usable for the FORTRAN 90 version of linefit. The usage is:

```
sav2data, '/irpolar/solar_mhd/yelles/6301_088200.sav'
```

This will create a directory structure similar to the `tar_prepare` routine described in Sect. 9.1.

A Installing FORTRAN 90

Download F90 compiler for Linux at <http://www.intel.com> → software → Compilers → Fortran Compilers for Linux → Free evaluation Download (direct link: <http://www.intel.com/software/products/compilers/flin/eval.htm>). Follow installation instructions.

B Installing MPICH

Download MPICH from <http://www-unix.mcs.anl.gov/mpi/mpich/>. Currently only MPICH-1 is supported (MPICH-2 support will be added on demand.)

In order to compile HELIX you have to set the environment variables for your f90 compiler, otherwise the f90 libraries are not created, eg:

```
export FC=/opt/intel/bin/ifort
export F90=/opt/intel/bin/ifort
```

Configure, make and install commands:

```
./configure --with-device=ch_p4 --prefix=/opt/mpich-1.2.5/
               --with-common-prefix=/opt/mpich-1.2.5/
make
make install
```

The prefix has to agree with the `MPI_DIR` in the `configure` script of your fortran directory.

Check `/opt/mpich-1.2.5/bin/mpif90` for the entries:

```
F90BASE="ifort"
F90LINKERBASE="ifort"
```

C Installing DISLIN

Download DISLIN from <http://www.dislin.de/> and follow installation instructions. If you plan to use MPICH and DISLIN you have to remove the `bzero` library from the DISLIN distribution (see note in Sect. 4.1).

D Running HELIX on Clusters

D.1 Running HELIX using MPICH at the MPS

When inverting several pixels the use of MPICH is highly recommended. The MPICH version will divide the list of pixels to several machines. MPICH has to be installed on the calling machine (from now on called the master). It is not necessary to have MPICH on the slaves (the machines which are used for the computation). See Section B.

MPICH can use several protocols to communicate with other machines. This description is based on the communication using `ssh`. You have to configure `ssh` for password-less authentication using a

public / private key pair. To test if password-less authentication is working, type the shell command in a terminal on the master:

```
ssh user@slave1 ls
```

This should list the directory content of the root directory on slave1. If you are asked for a password then read the ssh-documentation on how to switch to password-less authentication.

If the passwordless authentication works, but you are still unable to run HELIX using MPICH the reason might lie in your firewall setting. SuSE automatically activates the firewall, preventing `mpich` to work properly. Error messages similar to this one may appear:

```
p4_error: Timeout in making connection to remote process on lx40: 0
```

In this case try to disable the firewall completely or put your interface to the internal zone (SuSE - Yast - Firewall - Interfaces - Internal Zone).

The inversion must be run in a directory where the necessary files are accessible from the master and all slaves. You cannot use a local directory of one of the machines. The paths to the input-files (eg. profile archive) and the output files (eg. atmosphere archive) have to be the same for all machines (master and slaves). A good choice for a common directory is `/scratch/user` since this directory can be mounted from every machine at the MPS.

The machines used for MPICH are defined in a machine-file. Under Linux this file is `/opt/mpich-1.2.5/share/machines.LINUX` (assuming that `/opt/mpich-1.2.5/` is your MPICH installation path):

Every slave has to know the name of the master and all other slaves. Update your `/etc/hosts` file accordingly (or use the IP-numbers instead of the hostnames)!

```
# Change this file to contain the machines that you want to use
# to run MPI jobs on. The format is one host name per line, with either
#   hostname
# or
#   hostname:n
# where n is the number of processors in an SMP. The hostname should
# be the same as the result from the command "hostname"
lxlagg:2
lx40:4
lx41:4
lx42:4
lx43:4
seismo2:8
```

In this example the master is `lxlagg`, the slaves are `lxlagg`, `lx40` to `lx43` and `seismo2`.

The role of the master is only to distribute the pixels to the different slaves. This uses only very little computing power, therefore it makes sense to use the master also as slave (this is why `lagg-sxm2` is in the list twice). Use the `top` command on the different machines (master and slaves) to check the performance of MPICH.

Testing your MPICH environment

A small routine called `./bin/mpitest.$ARCH` is available to test the MPICH environment (`$ARCH` stands for the architecture you are using, for Linux-machines like `lx40` or `seismo2` it is `./bin/mpitest.mpich`, for helios it is `./bin/mpitest.solmpi` and for the gwdg-Linux Cluster it is `./bin/mpitest.scali`). If you want to test your setup on, e.g., `lx40` use the command:

```
lx40:/scratch/lagg/linefit> /opt/mpich/ch-p4/bin/mpirun \
    -machinefile ./machines.lx -np 9 bin/mpitest.mpich
```

This will produce the following output:

```
Hello, I am the master: lx40.mps.mpg.de
Hello, I am slave# 5 of 8 on lx40.mps.mpg.de
Hello, I am slave# 3 of 8 on lx43.mps.mpg.de
Hello, I am slave# 7 of 8 on lx42.mps.mpg.de
Hello, I am slave# 6 of 8 on lx41.mps.mpg.de
Hello, I am slave# 1 of 8 on lx41.mps.mpg.de
Hello, I am slave# 8 of 8 on lx43.mps.mpg.de
Hello, I am slave# 4 of 8 on seismo2.mps.mpg.de
Hello, I am slave# 2 of 8 on lx42.mps.mpg.de
```

Example 1

The command to run HELIX on 4 slaves (1 master + 4 slaves = 5 machines) is:

```
nice -15 /opt/mpich/ch-p4/bin/mpirun -np 5 bin/linefit.mpich -i mpitest.ipt
```

All slaves have to be accessible via ssh (without password) and all files have to be reachable from all machines under the same path. If this is the case the inversion should run and the progress will be displayed.

If you do not have write permissions to your MPICH installation directory you can use the option **-machinefile** to specify the file location. Example:

```
nice -15 /opt/mpich/ch-p4/bin/mpirun -np 5 \
    -machinefile ./machines.LINUX bin/linefit.mpich -i mpitest.ipt
```

The **nice -15** preceding the **mpirun** command gives the processes a lower priority. You should decrease the priority (i.e. increase the 'nice' number) when other users are also using the same machine.

Example 2

A very useful property of MPICH is that you can run programs in parallel on machines with different architectures. It is therefore possible to run an inversion on, let's say, Linux and Solaris machines at the same time.

Passwordless authentication has to be enabled between the server and the slaves (see above). The MPICH-binaries for the different architectures must exist.

Example: To run an inversion on the Linux machines lx?? and on the Solaris machine helios at the MPS you submit the command on an lx-machine:

```
nice -15 /opt/mpich/ch-p4/bin/mpirun -arch mpich -machinefile ./machines.lx \
    -np 4 -arch solmpi -machinefile machines.solaris -np 5 \
    ./bin/linefit.%a -i mpitest.ipt
```

This example runs the inversion defined in **test.ipt** on the master (the machine where you typed in the command) plus 3 slave nodes defined in the machinefile **machines.lx** and on 5 nodes of the helios cluster. The binaries **bin/linefit.mpich** and **bin/linefit.solmpi** must exist and be accessible from both architectures.

D.2 Running HELIX on the Solaris Cluster helios

The helios workstation is a Sun-Solaris workstation with currently 9 CPUs. An MPI-version of the code is available, the binary to be used is in the `bin` directory of your linefit distribution (`linefit.solmpi`). The solaris MPI-version does not support the ssh authentication described in Sect. D.1. It uses the less secure, but faster `rsh` protocol. Password-less rsh-authentication must be set up in order to use MPI on solaris: A file called `.rhosts` must exist in your home-directory. It must contain one line:

```
helios username
helios.mps.mpg.de username
```

(where "username" has to be replaced by your username). You can test the password-less rsh authentication by using the command

```
helios: > rsh helios.mps.mpg.de ls
```

This should give you a directory listing on helios without password query.

To run the MPI version on solaris use the commands in your linefit-directory:

```
. setup mpich
mpirun -np 10 bin/linefit.solmpi -i test.ipt
```

This should run the inversion on 9 CPUs plus a master process on CPU 0 (which uses almost no computing power).

For more advanced users it is possible to use the solaris cluster in combination with other Linux machines in house. Please read the documentation of mpirun.

D.3 Running HELIX on the Clusters gwdu102 and gwds1

The Linux Cluster in Göttingen (gwdu102) consists of 99 double processor Xeon 3.06GHz boards. It is ideally suited for inverting large maps. The MPI implementation in Göttingen is called SCALI.

Since January 2007 a new SGI Altix 4700 cluster exists: It has 512 CPU cores. Both clusters share the same home directory and temporary disk space.

Please read the homepages of the GWDG carefully before using the clusters:

<http://www.gwdg.de/service/index.html>. A description of the queuing system can be found here: http://www.gwdg.de/service/rechenanlagen/paralleltrechner/cluster_beschreibung/lsf.html.

D.4 Cluster gwdu102.gwdg.de

To compile the code for SCALI you first have to copy the linefit directory to the master node gwdu102.gwdg.de. The compile command is (in the `fortran` directory):

```
./configure scali
make linefit
```

You can also use the precompiled binary `./bin/linefit.scali`.

This should create a file `bin/linefit.scali` and a link to this binary in the linefit root directory. You should not use the X11 option. Although it should work, it does not make sense to display the data for 32 processes.

To run the job you use the gwdg-script `bsub` which calls `mpirun`:

```
bsub -u user@mps.mpg.de -q gwdg-pcpar -a scampi -n 16 -W 03:00 \
    ./bin/linefit.scali -i 13may01.018_2comp.ipt
```

This command submits the inversion to the lsf-queue using 16 processors (8 nodes) with a maximum time of 3 hours. After job completion an e-mail is sent to user@mps.mpg.de.

To control your job you can use the command:

bjobs

To delete your job use

bkill

To watch the CPU usage you can use the graphical tool

scadesktop

Make sure that your input file is correct and well tested! Also check if the directory structure is okay (eg. output directory `./atm_archive` must exist). Remember that the quotas on the home directory are quite low, use the `/scratch/user/` directory instead!

D.5 Cluster gwds1.gwdg.de

To compile the code for the SGI ALTix cluster you first have to copy the linefit directory to the master node gwds1.gwdg.de. The compile command is (in the `fortran` directory):

```
./configure sgialtix
make linefit
```

Note: There is no need to recompile the code. You can also use the precompiled binary

```
./bin/linefit.sgialtix.
```

You can test if your job runs successfully using the `mpirun` command. You should use `renice` the job to priority 19 and stop it after a few minutes by pressing CTRL-C. Running a job like this is only allowed for test purposes!

```
nice -19 mpirun -np 4 ./bin/linefit.sgialtix -i myinput.ipt
```

To submit the job to the queuing system use the gwdg-script `bsub`:

```
bsub -n 24 -W 05:00 -u user@mps.mpg.de -q gwdg-ia64 pam -mpi -auto_place \
    ./bin/linefit.sgialtix -i myinput.ipt
```

E Input File Examples

2-component atmosphere, test for convergence strategies File: `input/lmpik_test.ipt`

```
;----- COMMENTS -----
2-component fit
test for convergence strategy PIK_LM
;----- DIRECTORIES -----
PS          ./ps/          ;directory for postscript output
SAV         ./sav/         ;directory for storing results (sav-file)
```

```

PROFILE_ARCHIVE /data/gbso/VTT-data/data_may01/13may01/ ;input directory for
; profiles / observations
ATM_ARCHIVE ./atm_archive/ ;output directory for results (atmospheres)
ATM_SUFFIX lmpik_test_minfit90 ;add a suffix to the atm-directory to
; identify this run
WGT ./wgt/ ;directory for wgt-files
ATOM ./atom/ ;directory for atomic data files
;----- CONTROL -----
DISPLAY_MAP 0 ;if 1 then display maps of parameters after
; successful run (multiple pixels only!)
DISPLAY_PROFILE 0 ;if 1 then display fitted & observed profiles
DISPLAY_COMP 0 ;if 1 then display individual atm. components of
; fitted profiles
VERBOSE 0 ;verbosity of output: 0=quiet, 1=normal, 2=verbose
SAVE_FITPROF 0 ;if 1 then save fitted profile as sav-file (can be
; used as input profile)
OUTPUT X ;set to 'PS' for postscript output
;----- DATA SET -----
OBSERVATION 13may01.014cc ;observation in one of three formats: 1.) sav file
; of the format 'obsname'.profiles.sav, 2.
; spinor-profile, 3. sav-file created with linefit
; 'SAVE_FITPROF' keyword
WL_RANGE 10825.0000 10832.0000 ;WL-range to be used for analysis (may be
; only a part of the observation WL-range)
WL_NUM 256 ;# of WL-points (for synthesis only)
WL_DISP 0.0000 ;WL-calibration: dispersion per WL-bin number
WL_OFF 0.0000 ;WL-calibration: offset (used if != 0)
WL_BIN 1 ;wavelength binning
XPOS 0 147 ;two-elements vector containing xmin,xmax of the
; observation map to be analyzed
YPOS 0 96 ;two-elements vector containing ymin,ymax of the
; observation map to be analyzed
STEPX 1 ;step size for going from xmin to xmax
STEPLY 1 ;step size for going from ymin to ymax
AVERAGE 1 ;if 1 then average observation over the stepx/stepy
; size
SCANSIZE 0 ;stepsize of multiple scans within one observation
SYNTH 0 ;if 1 then create synthetic profile
NOISE 0.0000 ;noise level for adding artificial random noise
SMOOTH 0 0 ;smooth-value for profiles and smooth-method:
; (0=IDL-smooth function,1=FFT Low-Pass)
MEDIAN 3 ;median filter for observed profiles (0=off, >=2
; median filter width)
STRAYPOL_AMP 0.0000 ;amplitude for stray-polarization (only used for
; synthesis)
STRAYPOL_CORR 0 B ;iteration steps and orientation of
; scattering-polarization correction. Orientation:
; 'B' = along B-field, 'X' = a number defining an
; angle manually
SLIT_ORIENTATION 172.90 ;slit-orientation (used for straypol-corr)
SOLAR_POS 0.00 0.00 ;pos. of observation (x and y in arcseconds)
SOLAR_RADIUS 950.00 ;radius of sun in arcsec for time of obs.
;----- POST PROCESSING -----
CONV_FUNC ;convolution function of instrument
CONV_NWL 0 ;# of bins for convolution (used if number of
; WL_bins in data is small)
;----- ATMOSPHERES -----
NCOMP 2 ;number of components
; --- atmospheric component 1 ---
; NAME Value SCL_MIN SCL_MAX FIT %RG FIT
BFIEL 892.22 50.00 2000.00 4 ;magnetic field value in Gauss
AZIMU 37.05 -90.00 90.00 3 ;azimut of B-vector [deg]
GAMMA 43.50 0.00 180.00 2 ;inclination of B-vector [deg]
VELOS 1429.54 -7000.00 7000.00 1 ;line-of-sight velocity in m/s
VDAMP 0.01 0.01 0.70 0 ;damping constant (Voigt only)
VDOPP 0.42 0.10 1.00 1 ;doppler broadening (Voigt only)
EZERO 1.00 0.00 10.00 0 ;amplitude of components of
; propagation matrix (Voigt profile only!)
SGRAD 2.60 1.00 8.00 -5 ;gradient of source function
ALPHA 0.59 0.01 0.99 1 ;Filling factor for this component
; (only with multiple components)
USE_ATOM he1083.0.dat ;atomic data file(s) to be used for this component
; --- atmospheric component 2 ---
; NAME Value SCL_MIN SCL_MAX FIT %RG FIT
BFIEL 1984.00 50.00 2000.00 4
AZIMU 30.61 -90.00 90.00 3
GAMMA 47.42 0.00 180.00 2
VELOS 39876.10 5000.00 40000.00 1
VDAMP 0.01 0.01 0.70 0

```

```

VDOPP      0.99      0.10      1.00  1
EZERO      1.00      0.00      10.00 0
SGRAD      2.60      1.00      8.00 -5
ALPHA      0.41      0.01      0.99  1
USE_ATOM    he1083.0.dat
;----- LINE PARAMETERS -----
; --- fit parameters for line 1 ---
; NAME      Value      SCL_MIN  SCL_MAX  FIT
LINE_ID 10829.0911 ;wavelength to identify the line
LINE_STRENGTH 1.00      0.00      0.00  0 ;factor to adjust line strength
LINE_WLSHIFT 0.00      0.00      0.00  0 ;adjust central WL of line
; --- fit parameters for line 2 ---
; NAME      Value      SCL_MIN  SCL_MAX  FIT
LINE_ID 10830.2501 ;wavelength to identify the line
LINE_STRENGTH 1.00      0.00      0.00  0 ;factor to adjust line strength
LINE_WLSHIFT 0.00      0.00      0.00  0 ;adjust central WL of line
; --- fit parameters for line 3 ---
; NAME      Value      SCL_MIN  SCL_MAX  FIT
LINE_ID 10830.3397 ;wavelength to identify the line
LINE_STRENGTH 1.00      0.00      0.00  0 ;factor to adjust line strength
LINE_WLSHIFT 0.00      0.00      0.00  0 ;adjust central WL of line
;----- BLENDS -----
NBLEND      1 ;number of telluric blends
; --- telluric blend 1 ---
; NAME      Value      SCL_MIN  SCL_MAX  FIT
BLEND_WL      0.00      0.00      0.00  0 ;WL and WL-range for blend
BLEND_WIDTH    0.00      0.00      0.00  0 ;width of voigt-profile
BLEND_DAMP     0.00      0.00      0.00  0 ;damping of voigt-profile
BLEND_A0       0.00      0.00      0.00  0 ;amplitude of voigt-profile
; (blend is not used if BLEND_A0=0)
;----- GENERAL FIT PARAMETERS -----
; NAME      Value      SCL_MIN  SCL_MAX  FIT
CCORR        1.00      0.90      1.10  0 ;factor for continuum correction
STRAYLIGHT    0.00      0.00      1.00  0 ;straylight correction
; (non-dispersive, non-polarized)
;----- ANALYSIS METHOD -----
IQUV_WEIGHT  1.0000 1.0000 1.0000 1.0000 ;4-element vector defining relative
; weighting of IQUV (in that order). Additionally
; the code does an automatic weighting according to
; the strength of the I signal compared to the QUV
; signals. This weighting scheme is used in the
; PIKAIA fit routine.
WGT_FILE      he_default.wgt ;file with WL-dependent weighting function for IQUV
PROFILE        voigt ;functional form for pi- and sigma components of
; spectral line. Available: gauss, voigt or
; voigt_phys
MAGOPT        1 ;include magneto-optical effects (dispersion
; coefficients, (Voigt profile only!))
USE_GEFF       0 ;use effective Lande factor (=1) or real Zeeman
; pattern (=0)
USE_PB         1 ;if set, the zeeman-splitting and strength includes
; the Paschen-Back effect (from the table by
; Socas-Navarro)
PB_METHOD      poly ;use polynomials (=poly) or table interpolations
; (=table) to calculate the PB-effect
;----- PIKAIA PARAMATERS -----
CODE          FORTRAN ;PIKAIA code to use. Available: FORTRAN (=fast) or
; IDL (=platform independent).
METHOD        PIK_LM 1 20 0.90 ;minimization method: PIKAIA, POWELL (fast),
; LMDIF (fast) or PIK_LM (combined, for maps)
NCALLS        500 ;number of iterations in PIKAIA routine / max.
; number of calls for POWELL or LMDIF
;----- END OF INPUT FILE -----

```

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ODRPACK95 Version 2.01: Software for Weighted Orthogonal Distance Regression

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G Known Bugs

G.1 Normalization Problem (Jan-19 2007)

Until January 19th 2007 the code used a slightly wrong method to sum up the individual profiles resulting from different atomic lines / transitions. The calculated profiles were scaled with the reciprocal of the number of atomic lines / transitions (e.g. in the case of an inversion using the He 1083 nm line the absorption signature was weakened by a factor of 1/3).

This scaling was removed. Now every single line produces the same absorption signal, no matter on how many lines / transitions are calculated. The results from the old and the new runs are 1:1 exchangeable. All parameters calculated with the old method are still valid, only the parameter **SGRAD** will be affected: the value of **SGRAD** is decreased by a factor of one over the number of lines.

The effect of this error was that the **SGRAD** value changes when the number of atomic lines / transitions is changed. In the updated version **SGRAD** is independent on the number of lines (as it should be).

In order to maintain compatibility with old results the keyword **OLD_NORM** was introduced (for a description of the keyword see page 11). If this keyword is set to one, then the code uses the old normalization method, resulting in (number of lines)× higher values of **SGRAD**. Not setting this keyword means that the code uses the new (and hopefully correct) normalization.

and many others bugs... If you find a bug or an undesired result / behavior please describe it briefly and send it with all files necessary to reproduce the error (data, weighting file, input file, atomic data file) to the author.

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