

JURASSIC - scatter

Simulation Laboratory Climate Science (SimLab Climate),
Jülich Supercomputing Centre (JSC),
Forschungszentrum Jülich GmbH, Jülich, Germany ¹

28th March 2019

¹Contact: l.hoffmann@fz-juelich.de, s.griessbach@fz-juelich.de

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1 Introduction

The Juelich Rapid Spectral Simulation Code (JURASSIC) is a fast radiative transfer model for the mid-infrared spectral region (Hoffmann, 2006). It was used in several studies for the infrared limb sounder Michelson Interferometer for Passive Atmospheric Sounding (MIPAS) (Hoffmann et al., 2005, 2008), Cryogenic Infrared Spectrometers and Telescopes for the Atmosphere - New Frontiers (CRISTA-NF) (Hoffmann et al., 2009; Weigel et al., 2010), and Gimballed Limb Observer for Radiance Imaging of the Atmosphere (GLORIA) (Ungermann et al., 2010) and the nadir instrument Atmospheric Infrared Sounder (AIRS) (Hoffmann and Alexander, 2009; Grimsdell et al., 2010; Hoffmann et al., 2013).

For fast simulations, it applies pre-calculated look-up tables of spectral emissivities and approximations to radiative transfer calculations, namely the emissivity growth approximation (EGA) (Weinreb and Neuendorffer, 1973; Gordley and Russell, 1981; Marshall et al., 1994). The look-up-tables were calculated with the Reference Forward Model (RFM) (Dudhia et al., 2002; Dudhia, 2014, 2017), which is an exact line-by-line model specifically developed for MIPAS. For selected spectral windows, JURASSIC has been compared to the line-by-line models RFM and Karlsruhe Optimized and Precise Radiative transfer Algorithm (KOPRA) (Stiller, 2000; Stiller et al., 2002; Höpfner and Emde, 2005) and shows good agreement (Griessbach et al., 2013).

JURASSIC contains a scattering module that allows for radiative transfer simulations including single scattering on aerosol and cloud particles (Grießbach, 2012; Griessbach et al., 2013). Forward simulations with scattering on volcanic ash, ice and sulfate aerosol have been used to develop and characterize a volcanic ash detection method for MIPAS (Griessbach et al., 2012, 2014) and to discriminate between ice, ash, and sulfate aerosol (Griessbach et al., 2016, 2018).

Retrieval of large satellite data sets (e.g. AIRS) require plenty of computing time that can be provided by supercomputers. JURASSIC-scatter showed best performance using a hybrid MPI/openMP parallelization on the past JUROPA and JUQUEEN and the recent JURECA and JUWELS at the Jlich Supercomputing Centre (JSC), Forschungszentrum Jlich GmbH. It can also be used on workstations or laptops using either pure MPI or openMP parallelization.

This documentation is neither ready nor perfect, but it is a start to introduce you to JURASSIC(-scatter) and to enable you to work and to do science with this code package.

2 Getting Started

2.1 Quick Start

The following description is for Linux users. Download the code from github:

```
1 git clone https://github.com/slcs-jsc/jurassic-scatter.git
```

or as a .zip-file.

This will create the directory `jurassic-scatter` with the following subdirectories: `clim`, `docu`, `examples`, `lib`, `refrac`, `src`.

Go to `src`, open the Makefile and comment/uncomment the compiler sections, and modify the paths to the libraries according to your requirements. Then compile the code. If it finishes without error, go to `examples` run `clear.sh`, `aero0.sh` and `aero1.sh`. If this works you're done and you can start your own projects.

```
1 cd src
2 emacs Makefile
3 make
4 cd ../examples
5 clear.sh
6 aerosol0.sh
7 aerosol1.sh
```

If the makefile does not finish without any error either the path to the GSL is wrong or the library is not installed. Install the library and/or add the library and include paths to the makefile. If this does not work or if there are any other problems, please contact s.griessbach@fz-juelich.de or l.hoffmann@fz-juelich.de.

2.2 JURASSIC-scatter Repository

The JURASSIC-scatter repository contains 6 folders:

| folder | content |
|-----------------------|--|
| <code>clim</code> | climatology data from Remedios et al. (2007) |
| <code>docu</code> | this documentation |
| <code>examples</code> | example setups |
| <code>lib</code> | scripts to install the required GSL and NetCDF libraries (if required) |
| <code>refrac</code> | selection of pre-formatted refractive index files for scattering simulations |
| <code>src</code> | JURASSIC source files |

The JURASSIC source code is written in C and we usually compile the code with the GCC. The code only requires the GSL. In case the library is not installed and you do not have root access you can install the packages provided in `lib`.

2.3 How to do ...? - Examples

The following examples will show you how to use the JURASSIC modules, to do various forward simulations and retrievals. The detailed documentation of the JURASSIC modules is given in Section 5. For each example the required input files and output files are listed. The detailed descriptions of input and output files are given in Sections 3 and 4. The most substantial input file is the control file (Section 3.3). It contains all flags to setup the runs. We try to demonstrate the usage of all flags in the examples. The examples discussed here you will find in the `examples` folder as working examples.

2.3.1 Limb Clear Air Forward Simulation

Input files:

- `clear-air.ctl`: control file with global control flags
- `atm.tab`: atmospheric profile created for polar winter atmosphere
- `obs.tab`: observation geometry created for limb observer for 10 tangent altitudes
- `*.tab`: emissivity tables for each trace gas
- `*.filt`: filter functions for each spectral window

Output file:

- rad.tab: forward simulation results output file name

For this example we use the `climatology` and the `limb` module to create the required atmosphere and observation file, respectively. The forward simulation is started with the `formod` module.

```

1  #!/bin/bash
2
3  # path
4  src=../src
5
6  echo "Create atmosphere..."
7  $src/climatology clear-air.ct1 - atm.tab CLIMZONE pwin|| exit
8
9  echo "Create observation geometry..."
10 $src/limb clear-air.ct1 800 5 15 1 obs.tab || exit
11
12 echo "Call forward model..."
13 $src/formod clear-air.ct1 obs.tab atm.tab rad_clear.tab || exit

```

2.3.2 Nadir Aerosol Forward Simulation

Input files:

- aerosol0.ct1: control file with global control flags
- atm.tab: atmospheric profile created for polar winter atmosphere
- obs_nadir.tab: observation geometry created for nadir observer
- *.tab: emissivity tables for each trace gas
- *.filt: filter functions for each spectral window
- aero0.tab: aerosol/cloud geometry and microphysical properties
- complex refractive index file as given in aero.tab

Output file:

- rad_aero0.tab: forward simulation results output file name

For this example we use the `climatology` and the `nadir` module to create the required atmosphere and observation file, respectively. The forward simulation is started with the `formod` module.

```

1  #!/bin/bash
2
3  # path
4  src=./src
5
6  echo "Create atmosphere..."
7  $src/climatology aerosol1.ct1 - atm.tab CLIMZONE pwin || exit
8
9  echo "Create observation geometry..."
10 $src/nadir aerosol1.ct1 800 0 10 1 obs_nadir.tab || exit
11
12 echo "Call forward model..."
13
14 $src/formod aerosol0.ct1 obs_nadir.tab atm.tab rad_aero0.tab \
15 AEROFILE aero0.tab|| exit

```

2.3.3 Limb Aerosol Forward Simulation

Input files:

- aerosol1.ct1: control file with global control flags
- atm.tab: atmospheric profile created for polar winter atmosphere
- obs.tab: observation geometry created for nadir observer
- *.tab: emissivity tables for each trace gas
- *.filt: filter functions for each spectral window
- aero1.tab: aerosol/cloud geometry and microphysical properties (aero2.tab: using external optical properties)
- complex refractive index file as given in aero.tab

Output file:

- rad_aero1.tab: forward simulation results output file name

For this example we use the `climatology` and the `limb` module to create the required atmosphere and observation file, respectively. The forward simulation is started with the `formod` module.


```

1  #!/bin/bash
2
3  # path
4  src=../src
5
6  echo "Create atmosphere..."
7  $src/climatology aerosol1.ct1 - atm.tab CLIMZONE pwin || exit
8
9  echo "Create observation geometry..."
10 $src/nadir aerosol1.ct1 800 0 10 1 obs.tab || exit
11
12 echo "Call forward model..."
13
14 $src/formod aerosol1.ct1 obs.tab atm.tab rad_aero1.tab \
15     AEROFILE aero1.tab|| exit

```

2.3.4 Limb Clear Air Retrieval

- .ctl: control file with global control flags
- atm.tab: atmospheric profiles/mixing ratios
- obs.tab: observations and geometry
- rad.tab: forward simulation results output file name ???
- tables
- measurement data converted to obs.tab
- maybe one example for clear air nadir
- another example for clear air limb

TO DO

3 Input Files

3.1 Aerosol File

The aerosol file (`aero.tab`) contains a single 1D cloud scenario with the parameters given in Table 1. An aerosol scenario can be composed of multiple aerosol models. Each aerosol model contains data for a single scattering model in a single layer. The aerosol data must be given sorted starting with the highest layer and ending with the lowest layer. Please make sure that the layers, including their transition layers, do not overlap. Multiple cloud models can be superposed within one layer (e.g. multi-modal lognormal size distribution, different refractive indices, different scattering models). For this, please make sure that top and bottom altitudes are exactly the same. An example is given below.

Table 1: Columns of `aero.tab` file

| | | |
|-------|---|---|
| # \$1 | = | aerosol layer top altitude [km] |
| # \$2 | = | aerosol layer bottom altitude [km] |
| # \$3 | = | transition layer thickness [km] |
| # \$4 | = | source for optical properties |
| # \$5 | = | refractive index file |
| # \$6 | = | particle concentration of log-normal mode in cm^{-3} |
| # \$7 | = | median radius of log-normal mode in μm |
| # \$8 | = | width of log-normal mode |

Do:

| | | | | | | | | |
|---|------|------|------|-----|----------------------------------|-------|-------|-----|
| 1 | 14.0 | 13.0 | 0.01 | MIE | ../refrac/ice-266K-warren.dat | 0.032 | 3.6 | 1.6 |
| 2 | 14.0 | 13.0 | 0.01 | MIE | ../refrac/h2so4-215K-shettle.dat | 340.0 | 0.065 | 1.7 |
| 3 | 14.0 | 13.0 | 0.01 | MIE | ../refrac/h2so4-215K-shettle.dat | 5.0 | 0.49 | 1.3 |
| 4 | 12.0 | 10.0 | 0.01 | MIE | ../refrac/h2so4-215K-shettle.dat | 340.0 | 0.065 | 1.7 |
| 5 | 12.0 | 10.0 | 0.01 | MIE | ../refrac/h2so4-215K-shettle.dat | 5.0 | 0.49 | 1.3 |
| 6 | 9.0 | 8.0 | 0.01 | MIE | ../refrac/ice-266K-warren.dat | 0.032 | 3.6 | 1.6 |

Don't:

| | | | | | | | | |
|---|------|------|------|-----|----------------------------------|-------|-------|------|
| 1 | 14.0 | 13.0 | 0.01 | MIE | ../refrac/ice-266K-warren.dat | 0.032 | 3.6 | 1.6 |
| 2 | 14.0 | 13.0 | 0.01 | MIE | ../refrac/h2so4-215K-shettle.dat | 340.0 | 0.065 | 1.75 |
| 3 | 14.0 | 12.0 | 0.01 | MIE | ../refrac/h2so4-215K-shettle.dat | 5.0 | 0.49 | 1.3 |

Column 3, the transition layer thickness, defines the distance where the particle concentration decreases to zero. We highly recommend to use sharp cloud edges or small transition layers e.g. 0.01 km.

Column 4, the source for the optical properties can be

- Mie: internal Mie code with Gauss-Hermite integration
- Ext: external database, e.g. for non-spherical particles *not implemented yet*

Column 5 contains the path and name of an external file. For the Mie code complex refractive indices are required. The refractive index file is described in Section 3.7. For an external database of optical properties the file name must be given here. The optical properties database format is described in Section 3.6.

Columns 6–8, for Mie calculations we use mono-modal log-normal particle size distributions with the parameters (particle concentration, median radius and width) given in columns 6 to 8. If an external database is used columns 6 to 8 must be set to “0”.

The aerosol/cloud and scattering related control file parameters are listed in Table 3.

Sabines hint:

Generally transition layers are a nasty idea. They make physical interpretations much more difficult, e.g see Griessbach et al. (2013) and Höpfner et al. (2009) scenarios 4 and 5 for the effects of transition layer sampling. I strongly recommend to avoid large transition layers as long as possible.

Anyway, real clouds usually have sharp cloud edges (e.g. lifting condensation level).

3.2 Atmosphere File

The atmosphere file (`atm.tab`) is a XX+1 column list containing the temperature, pressure and volume mixing ratio profiles on a certain altitude grid. The number of total columns increases with the number of trace gases. The column order of the trace gases must be the same as the trace gas name order given in the control file (Section 3.3). The program `climatology` (Section 5.1.2) can be used to generate some default atmosphere files with an arbitrary altitude grid. The atmosphere related control file parameters are listed in Table 3.

The atmosphere file is the setup information for forward simulations. For retrievals it is the a priori information. Retrieval results are written in the same format (atmosphere structure) into a new file (e.g `atm_res.tab`). The retrieval output file names are hard-coded.

Table 2: Columns of *atm.tab* file

| | | |
|----------|---|--|
| # \$1 | = | time (seconds since 2000-01-01T00:00Z) |
| # \$2 | = | altitude [km] |
| # \$3 | = | longitude [deg] |
| # \$4 | = | latitude [deg] |
| # \$5 | = | pressure [hPa] |
| # \$6 | = | temperature [K] |
| # \$7 | = | CO2 volume mixing ratio |
| # \$8 | = | H2O volume mixing ratio |
| # \$9 | = | O3 volume mixing ratio |
| # \$NN | = | XX volume mixing ratio |
| # \$NN+1 | = | window 0: extinction [1/km] |
| # \$NN+2 | = | window 1: extinction [1/km] |

3.3 Control File

The control file (e.g. `clear.ct1`) contains all general setup parameters for simulation and retrieval runs. All mandatory and optional parameters are listed in Tables 3 and 4. Examples for clear air and aerosol simulation setups can be found in the `examples` folder.

3.3.1 Aerosol and Clouds

To include scattering on aerosol and cloud particles into the simulations up to 15 scattering models can be used. One scattering model can be a single mode of a multi-modal size distribution. To simulate a three modal size distribution three scattering models must be used (and placed at the same altitude in the aerosol input file). For each model individual refractive indices may be used. It is also possible to combine Mie-scattering and external databases in the same aerosol/cloud layer. For a clear air simulation the number of scattering model must be set to 0.

It is also possible to define an aerosol/cloud scenario and to neglect the scattering by setting the multiple scattering flag to 0. In this case you can choose if you want to use either the absorption or the extinction coefficient of the aerosol/cloud layers. (See Höpfner and Emde (2005) to learn which is best for a particular scenario.)

For a single scattering simulation the multiple scattering flag must be set to 1. If it is set to 2 or larger multiple scattering is simulated by also scattering the incoming rays that are calculated for the scattering source term.

Table 3: Control flags

| flag name | purpose | default | options |
|----------------------------------|---|----------|--|
| Emitter | | | |
| NG EMITTER[NG] | number of emitter emitter name (trace gas name) | 0 “ ” | 0–15 List of emitter names in Section 3.3.4 |
| Radiance channels | | | |
| ND NU[ND] | number of radiance channels central wave number of each channel in cm^{-1} | 0 “ ” | 0–50 range: 600–3000 cm^{-1} ??? |
| Spectral windows | | | |
| NW | number of spectral windows | 1 | 1–5 |
| WINDOW[ND] | window index of each channel | 0 | |
| Emissivity look-up tables | | | |
| TBLBASE | look-up table path and prefix | “-” | |
| Hydrostatic equilibrium | | | |
| HYDZ | reference height for hydrostatic pressure profile | -999 | -999: skip this option in km |
| Continua | | | |
| CTM_CO2 | CO2-continuum | 1 | 0: off; 1: on |
| CTM_H2O | H2O-continuum | 1 | 0: off; 1: on |
| CTM_N2 | N2-continuum | 1 | 0: off; 1: on |
| CTM_O2 | O2-continuum | 1 | 0: off; 1: on |
| Aerosol and Clouds | | | |
| SCA_N | number of scattering models | 0 | 0–15 0: clear air (no ext- inction, no scattering) |
| SCA_MULT | multiple scattering - number of recursions; to be used if SCA_N ≥ 1 | 1 | 0: extinction only 1: single scattering ≥ 2 : multiple scattering |
| SCA_EXT | extinction coefficient; to be used if SCA_MULT = 0 | beta_a | beta_e: β_e beta_a: β_a |
| Atmosphere/Climatology | | | |
| ZMIN | atmosphere bottom altitude | 0 | in km |
| ZMAX | atmosphere top altitude | 0 | in km |
| DZ | atmosphere vertical grid | 1 | in km |

Table 4: Control flags

| flag name | purpose | default | options |
|-----------------------------------|--|---------|--|
| Ray-tracing | | | |
| REFRAC | refraction in the atmosphere | 1 | 1: on; 0: off |
| RAYDS | maximum step lengths for ray-tracing | 10 | 10 km: suitable for limb 1 km: suitable for nadir |
| RAYDZ | maximum vertical component of step length | 1 | 1 km is reasonable |
| TRANS | transition layer sampling step | 0.02 | 0.01-0.1 km is reasonable |
| Interpolation of atmospheric data | | | |
| IP | interpolation method | 1 | 1: profile 2: satellite track 3: Lagrangian grid |
| CZ | influence length for vertical interpolation for IP=3 | 0 | in km |
| CX | influence length for horizontal interpolation for IP=3 | 0 | in km |
| Field of view | | | |
| FOV | field-of-view data file | “-” | path+filename |
| Retrieval interface | | | |
| RETP_ZMIN | minimum altitude for pressure retrieval | -999 | in km |
| RETP_ZMAX | maximum altitude for pressure retrieval | -999 | in km |
| RETT_ZMIN | minimum altitude for temperature retrieval | -999 | in km |
| RETT_ZMAX | maximum altitude for temperature retrieval | -999 | in km |
| RETP_ZMIN[NG] | minimum altitude for volume mixing ratio retrieval | -999 | in km |
| RETP_ZMAX[NG] | maximum altitude for volume mixing ratio retrieval | -999 | in km |
| RETK_ZMIN[NW] | minimum altitude for extinction retrieval | -999 | in km |
| RETK_ZMAX[NW] | maximum altitude for extinction retrieval | -999 | in km |
| Output flags | | | |
| WRITE_BBT | use brightness temperature instead of radiance | 0 | 0: no; 1: yes |
| WRITE_MATRIX | write matrix data | 0 | 0: no; 1: yes |

3.3.2 Continua

TO DO kurze Beschreibung, welches Schema?

3.3.3 Emissivity look-up Tables

Basename for table files and filter function files. Look-up tables are created in instrument specific spectral resolution/sampling. The look-up tables are described in Section 3.8.

3.3.4 Emitter

A list of supported trace gases is given in Table 5. The reading routine is not case sensitive.

3.3.5 Field of View

TO DO

3.3.6 Ray-tracing

The lengths of the line-of-sight segments to be integrated is either determined by RAYDS (total segment length) or RAYDZ (z-component of segment length). For limb scenarios RAYDS is most likely the limiting value and in nadir scenarios RAYDZ will be the limiting value. For limb-scenarios $\text{RAYDS} = 10 \text{ km}$ is reasonable. In nadir and sub-limb-scenarios a segment length of 10 km is too much to sample steep atmospheric gradients (temperature, trace gases). Especially for scattering simulations, where limb, sub-limb and nadir paths are calculated a reasonable combination of RAYDS and RAYDZ is required for fast and accurate simulations. From our experience $\text{RAYDS} = 10 \text{ km}$ and $\text{RAYDZ} = 0.1 - 1 \text{ km}$ offer a good trade off for accuracy and efficiency.

For cloud and aerosol simulations with a transition layer larger than 20 m the parameter TRANSS refines the sampling grid within the transition layer. The default value is $\text{TRANS} = 20 \text{ m}$. (Please see further comments on ray-tracing and transition layers in Section 5.2.6.)

Table 5: Trace Gases

| Emitter Name | Formula | Name |
|--------------|--|---|
| C2H2 | C ₂ H ₂ | acetylene |
| C2H6 | C ₂ H ₆ | ethane |
| CCl4 | CCl ₄ | carbon tetrachloride |
| CH4 | CH ₄ | methane |
| ClO | ClO | chlorine monoxide |
| ClONO2 | ClONO ₂ | chlorine nitrate |
| CO | CO | carbon monoxide |
| COF2 | COF ₂ | carbonyl fluoride |
| F11 | CCl ₃ F | trichlorofluoromethane |
| F12 | CCl ₂ F ₂ | dichlorodifluoromethane |
| F13 | CClF ₃ | chlorotrifluoromethane |
| F113 | C ₂ Cl ₃ F ₃ | trichlorotrifluoroethane |
| F114 | C ₂ Cl ₂ F ₄ | 1,2-dichlorotetrafluoroethane; cryofluorane |
| F14 | CF ₄ | tetrafluoromethane |
| F22 | CHClF ₂ | chlorodifluoromethane |
| H2O | H ₂ O | water |
| H2O2 | H ₂ O ₂ | hydrogen peroxide |
| HCN | HCN | hydrogen cyanide |
| HNO3 | HNO ₃ | nitric acid |
| HNO4 | HNO ₄ | peroxynitric acid |
| HOCl | HOCl | hypochlorous acid |
| N2O | N ₂ O | nitrous oxide |
| N2O5 | N ₂ O ₅ | dinitrogen pentoxide |
| NH3 | NH ₃ | ammonia |
| NO | NO | nitric oxide |
| NO2 | NO ₂ | nitrogen dioxide |
| O3 | O ₃ | ozone |
| OCS | OCS | carbonyl sulfide |
| PAN | CH ₃ COO ₂ NO ₂ | peroxyacetyl nitrate |
| SF6 | SF ₆ | sulfur hexafluoride |
| SO2 | SO ₂ | sulfur dioxide |

3.3.7 Spectral Windows

Each channel can be assigned to a window e.g. to have a constant extinction for all channels.

3.4 Directory List

The directory list file (e.g. `dirlist-aero.asc`) contains a list of directories e.g. for different cloud setups. One directory must be given per line. It can be used to run various scenarios one after another, or in parallel.

Table 6: Example for *dirlist-aero.asc* file

aerosol/sulfate aerosol/ash_small aerosol/basalt_small aerosol/andesite_small aerosol/ash_large aerosol/ice_large

3.5 Observation File

The observation file (`obs.tab`) is a multi-column list containing the geometry information (time, observer, view point and tangent point position) and the (measured) radiances (and/or transmittances) for each channel. (View point and tangent point differ, because of atmospheric refraction.) The number of columns increases with the number of channels. For retrievals this file contains the measurements. For forward simulations this file defines the viewing geometry. Columns 1 to 7 are mandatory. The forward simulation output is written in the same format (observation structure) into a new file specified when calling the forward model (e.g. `rad.tab`). The JURASSIC modules `limb` and `nadir` (Sections 5.1.7 and 5.1.8) can be used to create an observation geometry.

Table 7: Columns of *obs.tab* file

| | | |
|----------|---|---|
| # \$1 | = | time [seconds since 2000-01-01T00:00Z] |
| # \$2 | = | observer altitude [km] |
| # \$3 | = | observer longitude [deg] |
| # \$4 | = | observer latitude [deg] |
| # \$5 | = | view point altitude [km] |
| # \$6 | = | view point longitude [deg] |
| # \$7 | = | view point latitude [deg] |
| # \$8 | = | tangent point altitude [km] |
| # \$9 | = | tangent point longitude [deg] |
| # \$10 | = | tangent point latitude [deg] |
| # \$11 | = | channel 792: radiance [W/(m ² sr cm ⁻¹)] |
| # \$NN | = | channel XX: radiance [W/(m ² sr cm ⁻¹)] |
| # \$NN+1 | = | channel 792: transmittance |
| # \$NN+2 | = | channel XX: transmittance |

3.6 Optical Properties Database

External data bases containing the optical properties (β_e , β_s , and the phase function) for e.g. non-spherical particles can be used instead of the MIE code for spherical particles. **TO DO** describe input file format...

3.7 Refractive Index File

The refractive index file contains the wavenumber dependent complex refractive indices in the format given in Table 8. A formatted collection can be found in the folder **refrac**. One file contains the data set for one temperature (given in the file name). All refractive indices are taken from the HITRAN database (Rothman et al., 2009) aerosol compilation. References to the original sources are given there.

Table 8: *Columns of the refractive index files*

| | | |
|-------|---|-------------------|
| # \$1 | = | wavenumber [cm-1] |
| # \$2 | = | real part |
| # \$3 | = | imaginary part |

3.8 Tables

TO DO Describe me!!!

4 Output Files

rad.tab, same structure as observation file

Table 9: *Columns of `rad.tab` file*

| | | |
|----------|---|---|
| # \$1 | = | time [seconds since 2000-01-01T00:00Z] |
| # \$2 | = | observer altitude [km] |
| # \$3 | = | observer longitude [deg] |
| # \$4 | = | observer latitude [deg] |
| # \$5 | = | view point altitude [km] |
| # \$6 | = | view point longitude [deg] |
| # \$7 | = | view point latitude [deg] |
| # \$8 | = | tangent point altitude [km] |
| # \$9 | = | tangent point longitude [deg] |
| # \$10 | = | tangent point latitude [deg] |
| # \$11 | = | channel 792: radiance [W/(m ² sr cm ⁻¹)] |
| # \$NN | = | channel XX: radiance [W/(m ² sr cm ⁻¹)] |
| # \$NN+1 | = | channel 792: transmittance |
| # \$NN+2 | = | channel XX: transmittance |

5 JURASSIC Modules and Libraries

5.1 Modules

5.1.1 brightness

The `brightness` program calculates the brightness temperature in K for a given radiance in $\text{W}/(\text{m}^2 \text{ sr cm}^{-1})$ and wave number in cm^{-1} .

Example

```
1 ../src/brightness [radiance] [wavenumber]
2 ../src/brightness 0.0231451 960
```

5.1.2 climatology

The `climatology` program can be used to generate the atmosphere file (`atm.tab`). This program interpolates the chosen climatology from Remedios et al. (2007) onto the defined altitude grid and adds the trace gases in the same order as given in the control file. One can choose between polar winter (pwin), polar summer (psum), midlatitude night (midln) and equatorial night (equn) atmosphere (example line 3). The default is the midlatitude atmosphere (example line 2). The input grid can be taken from an already existing `atm.tab` file (example line 4). The grid points e.g. with 500 m vertical spacing will be taken from the file, but not the data! The control file parameters relevant for the climatology module can be found in Table 3 in the Emitter and Atmosphere/Climatology sections.

The Remedios climatology `clim_remedios.tab` is in the `clim` folder. Either copy the climatology into the working directory or set a link.

Examples

```
1 ../src/climatology [control file] [input grid] [output] [optional]
2 ../src/climatology clear-air.ct1 - atm.tab
3 ../src/climatology clear-air.ct1 - atm.tab CLIMZONE pwin
4 ../src/climatology clear-air.ct1 atm2.tab atm.tab
```

5.1.3 collect

TO DO

5.1.4 formod

The `formod` module is the program that starts a forward simulation. The minimum required input are the control file (Section 3.3), the observation file (Section 3.5), the atmosphere file (Section 3.2) and the name of an output file (Section 4).

Optional are `TASK`, `DIRLIST` (Section 3.4) and `AEROFILE` (Section 3.1). The option `TASK c` calculates the contribution of each specified trace gas separately. The output files are named `rad.tab.GAS` (e.g. `rad.tab.H2O`) and the format is the same as for the `rad.tab` output file in Section 4.

The option `DIRLIST` provides a file with directories. Each directory must contain an observation file, atmosphere file and optionally an aerosol file with the same name, but different content. The `DIRLIST` is a feature that is used to distribute large simulation/retrieval sets among multiple cores, e.g. on a supercomputer.

The option `AEROFILE` is required for scattering simulations. The `AEROFILE` contains the aerosol/cloud altitude information and the corresponding microphysical cloud/ aerosol properties.

Examples

```
1  ../src/formod [control file] [observation file] [atmosphere file] \  
2  [output]  
3  ../src/formod clear-air.ct1 obs.tab atm.tab rad.tab  
4  ../src/formod clear-air.ct1 obs.tab atm.tab rad.tab TASK c  
5  ../src/formod clear-air.ct1 obs.tab atm.tab rad.tab \  
6  DIRLIST dirlist-orbit.asc  
7  ../src/formod aerosol1.ct1 obs.tab atm.tab rad.tab AEROFILE aero.tab  
8  ../src/formod aerosol1.ct1 obs.tab atm.tab rad.tab \  
9  DIRLIST dirlist-aero.asc AEROFILE aero.tab
```

Instead of giving `TASK`, `DIRLIST` and `AEROFILE` as input, it can also be specified in the control file:

```
AEROFILE = aero.tab  
DIRLIST = dirlist-clim.asc  
TASK = c
```

5.1.5 interpolate

TO DO

5.1.6 kernel

TO DO

5.1.7 limb

The `limb` program creates an observation file (Section 3.5) for a specified limb geometry. In the example line 1 an observation file with 11 view point altitudes between 5 and 15 km is created.

Example

```
1 ../src/limb [control file] [observer altitude] \  
2           [min view point altitude] [max view point altitude] \  
3           [delta altitude] [output]  
4 ../src/limb aerosol1.ctl 800 5 15 1 obs.tab
```

5.1.8 nadir

The `nadir` program creates an observation file (Section 3.5) for the nadir geometry. In the example line 1 an observation file with nadir trace consisting of 11 points from 0 to 10 N is created.

Example

```
1 ../src/nadir [control file] [observer altitude] [min latitude] \  
2           [max latitude] [delta latitude] [output]  
3 ../src/nadir aerosol1.ctl 800 0 10 1 obs_nadir.tab
```

5.1.9 planck

The `planck` program returns the Planck radiance in $\text{W}/(\text{m}^2 \text{ sr cm}^{-1})$ for a given temperature in K and wavenumber in cm^{-1} .

Example

```
1 ../src/planck [temperature] [wavenumber]  
2 ../src/planck 270 960
```

5.1.10 raytrace

The **raytrace** programme can be used to perform the raytracing for a given geometry in a given atmosphere and to dump out the line of sight points and certain diagnostics into the files **raytrace.tab** and **los.NR**, where NR indicates the number of the line of sight. Further input options are: **LOSBASE**, **MASSBASE** and **AEROFIL**.

AEROFIL in example line 4 includes an aerosol file, so that the fine sampling around the aerosol/cloud edges is included into the raytracing. (see Section 5.2.6)

LOSBASE creates the output file **los.NR** if no other name is specified. It is switched on by default. In example line 6 the output file **liosf.0** is created. Example line 7 shows how to switch it off.

MASSBASE in example line 8 creates the output files **mass.numdens.0** and **mass.coldens.0**. **TO DO** what is in these files?

Examples

```
1  ../src/raytrace [control file] [observer file] [atmosphere file] \  
2  [options]  
3  ../src/raytrace clear-air.ctf obs_raytrace.tab atm.tab  
4  ../src/raytrace clear-air.ctf obs_raytrace.tab atm.tab \  
5  AEROFIL aero1.tab  
6  ../src/raytrace clear-air.ctf obs_raytrace.tab atm.tab LOSBASE liosf  
7  ../src/raytrace clear-air.ctf obs_raytrace.tab atm.tab LOSBASE -  
8  ../src/raytrace clear-air.ctf obs_raytrace.tab atm.tab MASSBASE mass
```

5.1.11 retrieval

TO DO

5.1.12 tab2bin

The **tab2bin** program can be used to convert the ASCII tables to binary format. Using binary format tables significantly reduces the file I/O time during e.g. **formod** runs. This program should be run on the architecture where the **JURASSIC** programmes using these tables will be executed. Please note, binary tables converted on **JUROPA** cannot be used on **JUQUEEN** and vice versa.

Example

```
1 ../src/tab2bin [control file]
2 ../src/tab2bin clear_air.ctl
```

5.2 Libraries

5.2.1 atmosphere.h/c

contains routines that deal with atmospheric data, atmospheric composition

5.2.2 continua.h/c

contains routines to calculate CO₂, H₂O, N₂ and O₂ continua

5.2.3 control.h/c

contains routines to read and scan the control file

5.2.4 forwardmodel.h/c

contains functions relevant to the forward model, radiative transfer calculation, table interpolation,...

5.2.5 jurassic.h

contains macros, constants (Appendix A.1), dimensions (Appendix A.2), and global struct definitions

5.2.6 lineofsight.h/c

contains functions that deal with raytracing and the line of sight Aerosol and cloud sampling: Done in raytracing. Extra los points about 5 m above and below cloud edges.

5.2.7 misc.h/c

contains routines that are used by several other programmes and routines

5.2.8 retrievalmodel.h/c

contains functions that deal with the retrieval

5.2.9 scatter.h/c

contains routines concerning scattering on aerosol and clouds; mie scattering is following Bohren and Huffman (1983) code; Gauss-Hermite scheme used for integration over log-normal size distribution; scattering of solar radiation included

5.3 Makefile

A Lists of Constants and Dimensions

A.1 Physical Constants

Table 10: Physical constants in `jurassic.h`.

| name | purpose | value |
|----------|---|---------------|
| C1 | First spectroscopic constant ($c_1 = 2hc^2$) [$\text{W}/(\text{m}^2 \text{ sr cm}^{-4})$] | 1.19104259e-8 |
| C2 | Second spectroscopic constant ($c_2 = hc/k$) [K/cm^{-1}] | 1.43877506 |
| P0 | Standard pressure [hPa] | 1013.25 |
| RE | Mean radius of Earth [km] | 6367.421 |
| ME | Mass of Earth [kg] | 5.976e24 |
| SCA_TSUN | Temperature of the sun [K] | 5780 |

A.2 Dimensions

Table 11: Dimensions in *control.h*.

| name | purpose | value |
|----------|---|------------------------|
| LEN | Maximum length of ASCII data lines | 5000 |
| MMAX | Maximum size of measurement vector | (NRMAX \times NDMAX) |
| NMAX | Maximum size of state vector | (NQMAX \times NPMAX) |
| NQMAX | Maximum number of quantities | (2+NGMAX+NWMAX) |
| NWMAX | Maximum number of spectral windows | 5 |
| NDMAX | Maximum number of radiance channels | 50 |
| NGMAX | Maximum number of emitters | 15 |
| NLOS | Maximum number of LOS points | 1000 |
| NPMAX | Maximum number of atmospheric data points | 1000 |
| NRMAX | Maximum number of ray paths | 1000 |
| NSHAPE | Maximum number of shape function grid points | 10000 |
| NFOV | Number of ray paths used for FOV calculations | 5 |
| TBLNPMAX | Maximum number of pressure levels in emissivity tables | 40 |
| TBLNSMAX | Maximum number of source function temperature levels | 1201 |
| TBLNTMAX | Maximum number of temperatures in emissivity tables | 30 |
| TBLNUMAX | Maximum number of column densities in emissivity tables | 320 |
| SCAMOD | Maximum number of scattering models | 15 |
| NLMAX | Maximum number of aerosol/cloud layers | 10 |
| NRAD | Number of points for Gauss-Hermite integration | 170 |
| NTHETA | Number of scattering angles (from 0 to 180 deg) | 181 |
| REFMAX | Maximum number of refractive indices | 5000 |

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