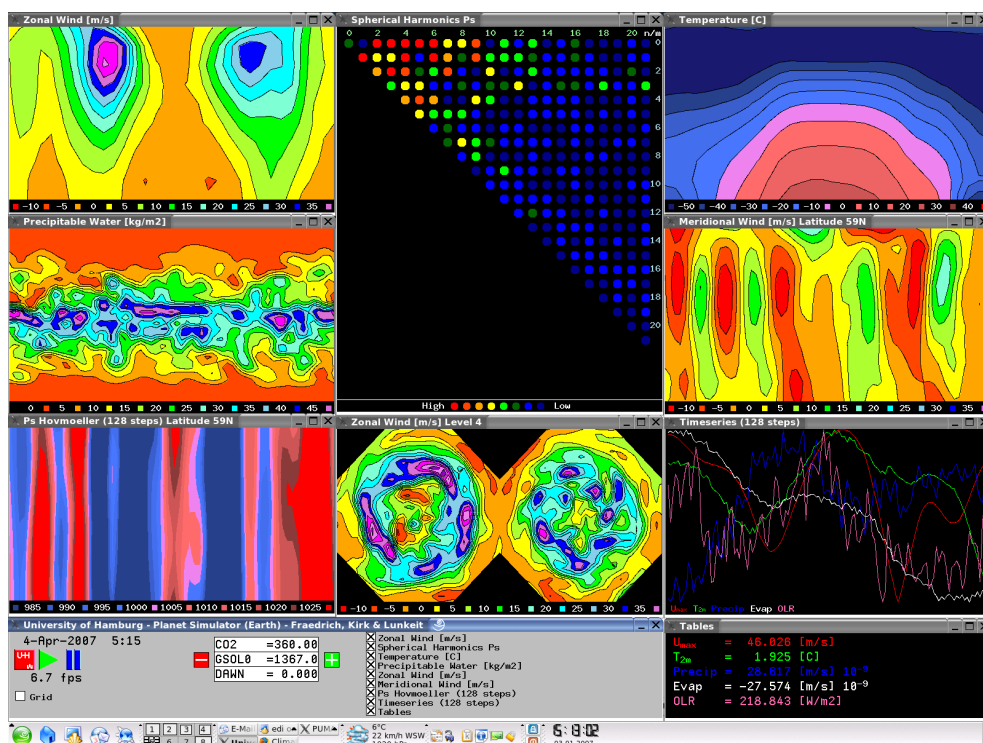




University of Hamburg

Planet Simulator



User's Guide

Version 16.0

Frank Lunkeit
Simon Blessing
Klaus Fraedrich
Heiko Jansen
Edilbert Kirk
Ute Luksch
Frank Sielmann

The Planet Simulator User's Guide is a publication of the Theoretical Meteorology at the Meteorological Institute of the University of Hamburg.

Address:

Prof. Dr. Klaus Fraedrich
Meteorological Institute
University of Hamburg
Bundesstrasse 55
D-20146 Hamburg

Contact:

Klaus.Fraedrich@zmaw.de
Frank.Lunkeit@zmaw.de
E.Kirk@gmx.de

Contents

1	Installation	5
1.1	Quick Installation	5
1.2	Most16 directory	5
1.3	Model build phase	6
1.4	Model run phase	7
1.5	Running long simulations	7
2	Modules	9
2.1	fluxmod.f90	10
2.2	miscmod.f90	11
2.3	surfmod.f90	12
2.4	fftmod.f90 / fft991mod.f90	13
2.5	landmod.f90	14
2.6	legmod.f90	16
2.7	mpimod.f90 / mpimod_stub.f90	17
2.8	outmod.f90	19
2.9	plasim.f90	20
2.10	plasimmod.f90	24
2.11	radmod.f90	25
2.12	rainmod.f90	27
2.13	seamod.f90	28
2.14	Sea ice and ocean modules	30
2.15	icemod.f90	33
2.16	oceanmod.f90	34
3	Parallel Program Execution	35
3.1	Concept	35
3.2	Parallelization in Gridpoint Domain	35
3.3	Parallelization in Spectral Domain	36
3.4	Synchronization points	36
3.5	Source code	37
4	Graphical User Interface	39
4.1	Graphical user interface (GUI)	39
4.2	GUI configuration	41
4.2.1	Array	42
4.2.2	Plot	43
4.2.3	Palette	43
4.2.4	Title	43
4.2.5	Geometry	43

5	Postprocessor Pumaburner	45
5.1	Introduction	45
5.2	Usage	45
5.3	Namelist	46
5.4	HTYPE	46
5.5	VTYPE	46
5.6	MODLEV	46
5.7	hPa	47
5.8	MEAN	47
5.9	Format of output data	47
5.10	SERVICE format	48
5.11	HHMM	48
5.12	HEAD7	48
5.13	MARS	48
5.14	MULTI	48
5.15	Namelist example	49
5.16	Troubleshooting	49
6	Graphics	51
6.1	Grads	51
6.2	Vis5D	54
7	Column Mode and Soundings	57
7.1	Setup	57
7.1.1	Basic switches for column setup	57
7.1.2	Boundary Conditions and forcing	57
7.2	Graphical User Interface (GUI)	58
A	List of Constants and Symbols	59
B	Planet Simulator Codes for Variables	63
C	Namelists	65
C.1	File puma_namelist	66
C.1.1	Namelist INP	66
C.1.2	Namelist PLANET	67
C.1.3	Namelist MISCPAR	67
C.1.4	Namelist FLUXPAR	68
C.1.5	Namelist RADPAR	68
C.1.6	Namelist RAINPAR	68
C.1.7	Namelist SURFPAR	69
C.2	File land_namelist	69
C.2.1	Namelist LANDPAR	69
C.3	File sea_namelist	70
C.3.1	Namelist SEAPAR	70
C.4	File ocean_namelist	70
C.4.1	Namelist OCEANPAR	70
C.5	File ice_namelist	70
C.5.1	Namelist ICEPAR	70

Chapter 1

Installation

The whole package containing the models "Planet Simulator" and "PUMA" along with "MoSt", the "Model Starter" comes in a single file, usually named "Most(n).tgz" with (n) specifying a version number. The following subsection gives an example, assuming version 16.

1.1 Quick Installation

```
tar -zxvf Most16.tgz
cd Most16
./configure.sh
./most.x
```

if your tar-command doesn't support the "-z" option (e.g. on Sun UNIX) type instead:

```
gunzip Most16.tgz
tar -xvf Most16.tar
cd Most16
./configure.sh
./most.x
```

If this sequence of commands produces error messages, consult the "FAQ" (Frequent Asked Questions) and README files in the **Most16** directory. They are plain text files, that can be read with the command "more" or any text editor.

1.2 Most16 directory

```
home/Most16> ls -lG
```

-rw-r--r--	1	1548	cc_check.c	<- Used by configure.sh
-rwxr-xr-x	1	57	cleanplasim	<- Delete run, bld and bin for PLASIM
-rwxr-xr-x	1	51	cleanpuma	<- Delete run, bld and bin for PUMA
drwxr-xr-x	2	4096	common	<- Topography files
-rwxr-xr-x	1	3911	configure.sh	<- The configure script
-rw-r--r--	1	308	csup.c	<- Currently unused
-rw-r--r--	1	234	f90check.f90	<- Used by configure.sh
-rw-r--r--	1	3033	FAQ	<- Frequently Asked Questions
drwxr-xr-x	2	4096	images	<- Directory for images
-rw-r--r--	1	154	makecheck	<- Used by configure.sh

```

-rw-r--r-- 1      85 makefile      <- Used to "make" most.x
-rw-r--r-- 1 107844 most.c         <- Source for MoSt (Model Starter)
-rw-r--r-- 1   6399 NEW_IN_VERSION_16 <- New in this version
drwxr-xr-x 8   4096 plasim        <- Planet Simulator directory
drwxr-xr-x 2   4096 postprocessor   <- Postprocessor directory
drwxr-xr-x 8   4096 puma          <- PUMA directory
-rw-r--r-- 1    839 README         <- Read this first
-rw-r--r-- 1   191 README_MAC_USER  <- Notes for MAC user
-rw-r--r-- 1   698 README_WINDOWS_USER <- Notes for Windows user

```

The directory structure must not be changed, even empty directories must be kept as they are, the Most program relies on the existence of these directories!

For each model, currently "Planet Simulator" and "PUMA" exists a directory (plasim) and (puma) with following subdirectories:

```
Most16/plasim> ls -lg
```

```

drwxr-xr-x 2   128 bin      <- model executables
drwxr-xr-x 2  1824 bld      <- build directory
drwxr-xr-x 2   280 dat      <- initial and boundary data
drwxr-xr-x 2    80 doc      <- documentation, user's guide, reference manual
drwxr-xr-x 2   928 run      <- run directory
drwxr-xr-x 2  1744 src      <- source code

```

After installation only "dat", "doc" and "src" contain files, all other directories are empty.

Running "Most" to setup a model configuration and define an experiment uses the directories in the following manner:

1.3 Model build phase

Most writes an executable shell script to the "bld" directory and executes it directly hereafter. It copies all necessary source files from "src" to "bld" and modifies them according to the selected parameter configuration. Modification of source code is necessary for vertical and horizontal resolution change and for using more than 1 processor (parallel program execution). The original files in the "src" directory are not changed by Most. The program modules are then compiled and linked using the "make" command (in bld/most_plasim_build), also issued by Most. Most provides a makefile named make_plasim for building the executable. For modules that exist in more than one version the selection of the module to use is done by environment variables that are set automatically by MoSt but may be changed manually by the user. Look into the make_plasim for further information. The resolution and CPU parameters are coded into the filename of the executable, in order to have different names for different versions. E.g. the executable "most_plasim.t21_l10_p2.x" is an executable compiled for a horizontal resolution of T21, a vertical resolution of 10 levels and 2 CPU's. The executable is copied to the models "bin" directory after building. Each time Most is used to setup a new experiment it checks the "bin" directory for a matching executable. If it's there, it's used without rebuilding otherwise a new executable with the selected parameters is created. Rebuilding may be forced by using the cleanplasim command in the Most directory. The build directory is not cleared after usage. The user may want to modify the makefile or the build script for his own purposes and start the building directly by executing "most_plasim_build". For permanent user modifications the contents of the "bld" directories have to be copied elsewhere, because each usage of Most overwrites the contents of "bld".

1.4 Model run phase

After building the model with the selected configuration, Most writes or copies all necessary files to the model's "run" directory. These are the executable, initial and boundary data, namelist files containing the parameter and finally the run script itself. Depending on the exit from Most, either "Save & Exit" or "Run & Exit", the run script is started from Most and takes control of the model run. A checkmark on GUI invokes also the Graphical User Interface for user control and display of variables during the run. Again all contents of the "run" directory are subject of change for the user. But it would be wise to keep changed run setups in other, user created directories, because each usage of Most overwrites the contents of the run directory.

1.5 Running long simulations

For long simulations make a new directory on a filesystem, that has enough free disk space to store the results. You may use the "df" command to check filesystems.

Hint 1: Don't use your home directory if there are filequotas. Your run may crash due to file quota exceeded.

Hint 2: Use a local disk, not NFS mounted filesystems if possible. The model runs much faster writing output to local disks.

Example:

- cd **Most16**
- ./most.x
- Select model and resolution
- Switch GUI off
- Switch Output on
- Edit number of years to run
- Click on "Save & Exit"
- Make a directory, e.g. mkdir /data/longsim
- cp plasim/run/* /data/longsim
- cd /data/longsim
- edit most_plasim_run for experiment name
- edit namelist files if necessary
- start simulation with most_plasim_run &

Chapter 2

Modules

In the following, the purposes of the individual modules is given and the general structure and possible input and output opportunities (namelist and files) are explained.

2.1 fluxmod.f90

General The module `fluxmod.f90` contains subroutines to compute the different surface fluxes and to perform the vertical diffusion. The interface to the main PUMA module `puma.f90` is given by the subroutines `fluxini`, `fluxstep` and `fluxstop` which are called in `puma.f90` from the subroutines `prolog`, `gridpointd` and `epilog`, respectively.

Input/Output `fluxmod.f90` does not use any extra input file or output file and is controlled by the namelist `fluxpar` which is part of the namelist file `puma_namelist`:

Parameter	Type	Purpose	Default
NEVAP	Integer	Switch for surface evaporation (0 = off, 1 = on)	1
NSHFL	Integer	Switch for surface sensible heat flux (0 = off, 1 = on)	1
NSTRESS	Integer	Switch for surface wind stress (0 = off, 1 = on)	1
NTSA	Integer	Switch for computing the near surface air temperature which is used for the Richardson number (1 = potential temperature, 2 = virtual potential temperature)	2
NVDIFF	Integer	Switch for vertical diffusion (0 = off, 1 = on)	1
VDIFF_LAMM	Real	Tuning parameter for vertical diffusion	160.
VDIFF_B	Real	Tuning parameter for vertical diffusion	5.
VDIFF_C	Real	Tuning parameter for vertical diffusion	5.
VDIFF_D	Real	Tuning parameter for vertical diffusion	5.
ZUMIN_D	Real	Minimum surface wind speed (m/s)	1.

Structure Internally, `fluxmod.f90` uses the FORTRAN-90 module `fluxmod`, which uses the global common module `pumamod` from `pumamod.f90`. Subroutine `fluxini` reads the namelist and, if the parallel version is used, distributes the namelist parameters to the different processes. Subroutine `fluxstep` calls the subroutine `surflx` to compute the surface fluxes and calls the subroutine `vdiff` to do the vertical diffusion. Subroutine `fluxstop` is a dummy subroutine since there is nothing to do to finalize the computations in `fluxmod.f90`. The computation of the surface fluxes in `surflx` is spitted into several parts. After initializing the stability dependent transfer coefficients, the subroutines `mkstress`, `mkshfl` and `mkevap` do the computations which are related to the surface wind stress, the surface sensible heat flux and the surface evaporation, respectively.

2.2 miscmod.f90

General The module `miscmod.f90` contains miscellaneous subroutines which do not fit well to other modules. The interface to the main module `plsim.f90` is given by the subroutines `miscini`, `miscstep` and `miscstop` which are called in `puma.f90` from the subroutines `prolog`, `gridpointd` and `epilog`, respectively. A subroutine to eliminate spurious negative humidity and an optional subroutine to relax the upper level temperature towards a prescribed distribution is included in `miscmod.f90`.

Input/Output `miscmod.f90` does not use any extra output file. If the relaxation is switched on, a climatological annual cycle of the prescribed upper level temperature distribution [K] is read from the external file `surface.txt`. The file format is formatted SERVICE format with (8I10) for the headers and (8E12.6) for the temperature fields. To assign the field, the header needs to have the header information code 130, level 1 and a date identifier of the form `yymmdd` or `mmdd` where `mm` goes from 1 to 12 (January to December) or from 0 to 14 (including the December of the previous year and the January of the following year). Fields which are not needed will be skipped. The module is controlled by the namelist `miscpar` which is part of the namelist file `puma_namelist`:

Parameter	Type	Purpose	default
NFIXER	Integer	Switch for correction of negative moisture (0 = off , 1= on)	1
NUDGE	Integer	Switch for temperature relaxation in the uppermost level (0 = off , 1= on)	0
TNUDGE	Real	Time scale [d] of the temperature relaxation	10.

Structure Internally, `miscmod.f90` uses the FORTRAN-90 module `miscmod`, which uses the global common module `pumamod` from `pumamod.f90`. Subroutine `miscini` reads the namelist and, if the parallel version is used, distributes the namelist parameters to the different processes. If the relaxation is switched on, the climatological temperature is read from `surface.txt` and distributed to the processors. Subroutine `miscstep` calls the subroutine `fixer` to eliminate spurious negative humidity arising from the spectral method and, if the relaxation is switched on, calls the subroutine `mknudge` to do the temperature nudging. Subroutine `miscstop` is a dummy subroutine since there is nothing to do to finalize the computations in `miscmod.f90`.

2.3 surfmod.f90

General The module `surfmod.f90` deals as an interface between the atmospheric part of the model and modules, or models, for the land and the oceans. The interface to the main PUMA module `puma.f90` is given by the subroutines `surfini`, `surfstep` and `surfstop` which are called in `puma.f90` from the subroutines `prolog`, `gridpointd` and `epilog`, respectively. Calls to subroutines named `landini`, `landstep` and `landstop` and `seaini`, `seastep` and `seastop` provide the interface to land and the ocean modules, respectively.

Input/Output `surfmod.f90` reads the land-sea mask and the orography (surface geopotential) [m^2/s^2] from file `surface.txt`. The file format is formatted SERVICE format with (8I10) for the headers and (8E12.6) for the fields. To assign the fields, the headers need to have the header information code 129 for the surface geopotential and 172 for the land-sea mask (1.0 = land; 0.0 = sea). Fractional land-sea-masks containing other values than 1.0 and 0.0 will be converted with values > 0.5 set to 1.0 and all other to 0.0. `surfmod.f90` is controlled by the namelist `surfpar` which is part of the namelist file `puma_namelist`:

Parameter	Type	Purpose	default
NSURF	Integer	Debug switch	not active
NOROMAX	Integer	Resolution of orography	NTRU
OROSCALE	Real	Scaling factor for orography	1.0

Structure Internally, `surfmod.f90` uses the FORTRAN-90 module `surfmod`, which uses the global common module `pumamod` from `pumamod.f90`. Subroutine `surfini` reads the namelist and, if the parallel version is used, distributes the namelist parameters to the different processes. If the run is not started from a restart file, the land-sea-mask and the orography are read from file `surface.txt`. According to the namelist input, the orography is scaled by `OROSCALE`, transferred into spectral space and truncated to `NOROMAX`. Calls to subroutines `landini` and `seaini` are the interfaces to the respective initialization routines contained in the land and ocean modules. During the run, the interface to land and ocean is given by calls to the external subroutines `landstep` and `seastep`, which are called by `surfstep`. At the end of the integration, interface subroutines `landstop` and `seastop` are called by `surfstop`.

2.4 `fftmod.f90` / `fft991mod.f90`

General The module `fftmod.f90` contains all subroutines necessary to perform the fast fourier transformation and its inverse. The interface to the main module `plasim.f90` is given by the subroutines `gp2fc` and `fc2gp` which are called in `plasim.f90` from the subroutine `gridpoint`.

Input/Output `fftmod.f90` does not use any extra input file or output file. No namelist input is required.

Structure Internally, `fftmod.f90` uses the FORTRAN-90 module `fftmod`, which uses no other modules. Subroutine `gp2fc` performs the transformation from grid point space into fourier space while the subroutine `fc2gp` does the transformation from fourier space into grid point space. Both routines use several subroutines to do the direct or indirect transformation for different factors. When `gp2fc` or `fc2gp` is called for the first time, `fftini` is called to do the initialization of the FFT.

The alternate module `fft991mod.f90` may be used instead of `fftmod.f90`. While `fftmod.f90` runs faster `fft991mod.f90` can be used for resolutions, that are not supported by `fftmod.f90`, e.g. T63 or T106. Edit the file `Most16/plasim/src/make_plasim` for module selection. Use either

```
FFTMOD=fftmod
```

or

```
FFTMOD=fft991mod
```

2.5 landmod.f90

General The module `landmod.f90` contains parameterizations for land surface and soil processes which include the simple biome model SIMBA and a model for the river runoff. The interface to the **Planet Simulator** is given via the module `surfmod.f90` by the subroutines `landini`, `landstep` and `landstop` which are called in `surfmod.f90` from the subroutines `surfini`, `surfstep` and `surfstop`, respectively.

Input/Output `landmod.f90` reads several surface and soil parameters either from the initial file `surface.txt` or from the restart file `plasim_restart` which is written at the end of an integration. `surface.txt` contains several surface fields which are needed for initialization. The file format is formatted SERVICE format with (8I10) for the header and (8E12.6) for the fields. The file may include the following fields: surface geopotential (orography) [m^2/s^2], land-sea mask [1.0,0.0], surface roughness [m], background albedo [frac.], glacier mask [frac.], bucket size [m], soil temperature [K], climatological annual cycle of the surface temperature [K], climatological annual cycle of the soil wetness [m]. To assign the fields, the headers need to have the header information code 129 for surface geopotential, code 172 for the land-sea mask (1. = land; 0. = sea), 173 for the surface roughness, 174 for the background albedo, 232 for the glacier mask (1. = glacier; 0. = no glacier), 229 for the bucket size, 209 for the soil temperature, 169 for the surface temperature and 140 for the soil wetness. for the climatological annual cycles of surface temperature and soil wetness, a date identifier of the form `yymmdd` or `mmdd` where `mm` goes from 1 to 12 (January to December) is required. Two additional months with `mm=0` indicating the December of the preceding year and `mm=13` for the January of the following year may be included for interpolation during transient simulations. If there are some fields not present in the `surface.txt` default values will be used which can be set in the namelist. The use of some fields depend on the setting of some namelist parameters. The restart file `plasim_restart` is an unformatted file which contains all variables needed to continue the run. `landmod.f90` is controlled by the namelist `landpar` given in the namelist file `land_namelist`:

Parameter	Type	Purpose	Default
NLANDT	Integer	Switch for surface temperature (1 = computed; 0 = climatology)	1
NLANDW	Integer	Switch for soil wetness (1 = computed; 0 = climatology)	1
NBIOME	Integer	Switch for biome model SIMBA (1 = on ; 0 = off)	0
ALBLAND	Real	Background albedo	0.2
DZ0LAND	Real	Roughnesslength [m]	2.0
DRHSLAND	Real	Wetness factor	0.25
ALBSMIN	Real	Minimum albedo for snow	0.4
ALBSMAX	Real	Maximum albedo for snow	0.8

Parameter	Type	Purpose	Default
NWATCINI	Integer	Switch to initialize soil water content manually (1 = on;0 = off)	0
DWATCINI	Real	Soil water content (m) for manual initialization	0.0
ALBGMIN	Real	Minimum albedo for glaciers	0.6
ALBGMAX	Real	Maximum albedo for glaciers	0.8
WSMAX	Real	Maximum field capacity of soil water (bucket size) [m]	0.5
DRHSFULL	Real	Threshold above which wetness factor is 1	0.4
DZGLAC	Real	Threshold of orography to be glacier (-1.0 = none) [m]	-1.0
DZTOP	Real	Thickness of the uppermost soil layer [m]	0.2
DSOILZ(5)	Real Array	Soil layer thicknesses [m]	0.4,0.8,1.6,3.2,6.4

Structure Internally, `landmod.f90` uses the FORTRAN-90 module `landmod`, which uses the global common module `pumamod` from `plasimmod.f90`. Subroutine `landini` reads the namelist and, if the parallel version is used, distributes the namelist parameters to the different processes. If the run is not started from a restart file, the initialization file `surface.txt` is being read. The soil and the river runoff are initialized via `soilini` and `roffini` and different variables are set according to the values given by the namelist or the `surface.txt`. Additionally, the climatological surface temperatures and soil wetnesses are updated from `surface.txt` if `NRESTART = 2`. If `NRESTART = 3` (special application) the bucket size, the roughness length and the albedo are set to the values given in the namelist. Subroutine `landstep` computes new surface and soil values via `soilstep` which calls `tands` and `wandr` for the heat and water budgets, respectively. If `NLANDT` and/or `NLANDW` are set to 0, climatological values are used for the surface temperature and the soil wetness. Via `roffstep` the river runoff is computed. Finally the biome model `simbastep` is called. The land model is finalized by `landstop` which writes the restart record to `plasim_restart`.

2.6 legmod.f90

General The module `legmod.f90` contains all subroutines necessary to perform the Legendre transformation and its inverse. The interface to the main module `plasim.f90` is given by the subroutines `legini`, `inigau`, `fc2sp`, `fc3sp`, and `sp2gp` which are called in `plasim.f90` from the subroutines `prolog` and `gridpoint`

Input/Output `legmod.f90` does not use any extra input file or output file. No namelist input is required

The following subroutines are included in `legmod.f90`:

Subroutine	Purpose
<code>inigau</code>	compute Gaussian abscissas and weights
<code>legini</code>	compute Legendre polynomials
<code>fs2sp</code>	Fourier to Spectral transformation
<code>sp2fc</code>	Spectral to Fourier transformation
<code>sp3fc</code>	Simultaneous transformation of T, Div., and Vort.
<code>dirlega</code>	Compute and transform adiabatic tendencies
<code>dirlegd</code>	Compute and transform diabatic tendencies
<code>invlega</code>	Spectral to Fourier - adiabatic part
<code>invlegd</code>	Spectral to Fourier - diabatic part

2.7 mpimod.f90 / mpimod_stub.f90

General The module `mpimod.f90` contains interface subroutines to the MPI (Message Passing Interface) needed for (massive) parallel computing. Several MPI routines are called from the module. The interface to other modules are given by numerous subroutines which names starts with *mp*. Subroutines from `mpimod.f90` are called in several other modules. There are no direct calls to MPI other than in `mpimod.f90`. This encapsulation makes it possible to use `mpimod_stub.f90` for single CPU runs without changing any other part of the model code. The selection is done automatically by using MoSt or manually by editing "Most15/plasim/src/make_plasim".

Input/Output `mpimod.f90` and `mpimod_stub` do not use any extra input file or output file. No namelist input is required

Structure Internally, `mpimod.f90` uses the FORTRAN-90 module `mpimod`, which uses the global common module `pumamod` from `plasimmod.f90` and the MPI module `mpi`. The following subroutines are included in `mpimod.f90`:

Subroutine	Purpose
<i>mpbci</i>	broadcast 1 integer
<i>mpbcin</i>	broadcast n integers
<i>mpbcr</i>	broadcast 1 real
<i>mpbcrn</i>	broadcast n reals
<i>mpbcl</i>	broadcast 1 logical
<i>mpscin</i>	scatter n integers
<i>mpscrn</i>	scatter n reals
<i>mpscgp</i>	scatter grid point field
<i>mpgagp</i>	gather grid point field
<i>mpgallgp</i>	gather grid point field to all
<i>mpscsp</i>	scatter spectral field
<i>mpgasp</i>	gather spectral field
<i>mpgacs</i>	gather cross section
<i>mpgallsp</i>	gather spectral field to all
<i>mpsum</i>	sum spectral field
<i>mpsumsc</i>	sum and scatter spectral field
<i>mpsumr</i>	sum n reals
<i>mpsumbcr</i>	sum and broadcast n reals
<i>mpstart</i>	initialize MPI
<i>mpstop</i>	finalize MPI

Subroutine	Purpose
<i>mpreadgp</i>	read and scatter grid point field
<i>mpwritegp</i>	gather and write grid point field
<i>mpwritegph</i>	gather and write (with header) grid point field
<i>mpreadsp</i>	read and scatter spectral field
<i>mpwritesp</i>	gather and write spectral field
<i>mpi_info</i>	give information about setup
<i>mpgetsp</i>	read spectral array from restart file
<i>mpgetgp</i>	read gridpoint array from restart file
<i>mpputsp</i>	write spectral array to restart file
<i>mpputgp</i>	write gridpoint array to restart file
<i>mpmaxval</i>	compute maximum value of an array
<i>mpsumval</i>	compute sum of all array elements

2.8 outmod.f90

General The module `outmod.f90` controls the data output of the model. The interface to the main PUMA module `puma.f90` is given by the subroutines `outini`, `outgp`, `outsp`, `oureset` and `outaccu` which are called in `puma.f90` from the subroutines `prolog` and `master`.

Input/Output `outmod.f90` writes the output data to the file `puma_output` which is an unformatted file. `puma_output` is designed to be post processed by the program `burn` (see section 5), which converts the model variables to useful output in user friendly format. There is no separate namelist for `outmod.f90` but some parameter of namelist `inp` of `plasim.f90` are used to control the format and the output interval.

Structure Internally, `outmod.f90` uses the global common module `pumamod` from `plasimmod.f90` in several subroutines. Subroutine `outini` does the initialization. Subroutines `outgp` and `outsp` write the grid point and the spectral fields to the output file `puma_output`. `outaccu` accumulates some output variables over the output interval. `oureset` resets the accumulated arrays to zero.

2.9 plasim.f90

General The module `plasim.f90` is the main module of the model. It includes the main program `plasim` and controls the run. From `plasim.f90` the interface routines to the modules `miscmod.f90`, `fluxmod.f90`, `radmod.f90`, `rainmod.f90`, `surfmod.f90` are called. The output is done by calling the interface routines to `outmod.f90`. In addition, the adiabatic tendencies and the horizontal diffusion are computed in `plasim.f90`. To do the necessary transformations, calls to the modules `fftmod.f90` and `legmod.f90` are used.

Input/Output `plasim.f90` does not use any extra input file or output file. A diagnostic print out is written on standard output. `plasim.f90` is controlled by the namelist `inp` which is part of the namelist file `puma_namelist`:

Parameter	Type	Purpose	Default
COLUMN	Integer	1: Set all parameters for default column mode	0
KICK	Integer	Switch for initial white noise disturbance on surface pressure (0 = none; 1 = global; 2 = hemispherically symmetric; 3 = one wavenumber only)	1
NWPD	Integer	Number of Writes Per Day (for output data)	1
NADV	Integer	Switch for advection (0 = off; 1 = on)	1
NCOEFF	Integer	Number of spectral coefficients in diagnostic print out	0
NDEL(NLEV)	Integer Array	Order of the horizontal diffusion	NLEV · 2
NDIAG	Integer	Time interval for diagnostic print out [time steps]	12
NKITS	Integer	Number of initial explicit Euler time steps	3
N_RUN_YEARS	Integer	Number of years to run	1
N_RUN_MONTHS	Integer	Number of months to run	0
N_RUN_DAYS	Integer	Number of days to run (for short test runs)	-1
N_START_YEAR	Integer	Start year	1
N_START_MONTH	Integer	Start month	1
N_DAYS_PER_YEAR	Integer	365: use real calendar with leap years, 360: use simple calendar with 12 months of equal length	360
N_DAYS_PER_MONTH	Integer	Number of days per month for simple calendar	30

Parameter	Type	Purpose	Default
MPSTEP	Integer	Minutes per step = length of timestep	45
NEQSIG	Integer	Switch for non equally spaced sigma levels (1 = non equally spaced; 1 = equally spaced)	1
NPRINT	Integer	Switch for extended debug print out (0 = off; 1 = on; 2 = very extended)	0
NPRHOR	Integer	Number of the grid point to be used for very extended debug print out	0
NPACKSP	Integer	Switch for spectral output (0 = normal; 1 = compressed)	1
NPACKGP	Integer	Switch for grid point output (0 = normal; 1 = compressed)	1
NRAD	Integer	Switch for radiation (0 = off; 1 = on)	1
NFLUX	Integer	Switch for surface fluxes and vertical diffusion (0 = off; 1 = on)	1
NDIAGGP	Integer	Switch for additional diagnostic grid point output (0 = off; 1 = on)	0
NDIAGSP	Integer	Switch for additional diagnostic spectral output (0 = off; 1 = on)	0
NDIAGCF	Integer	Switch for additional cloud forcing diagnostic (0 = off; 1 = on)	0
NDIAGGP2D	Integer	Number of additional diagnostic 2-d grid point output (0 = off; 1 = on)	0
NDIAGGP3D	Integer	Number of additional diagnostic 3-d grid point output (0 = off; 1 = on)	0
NDIAGSP2D	Integer	Number of additional diagnostic 2-d spectral output (0 = off; 1 = on)	0
NDIAGSP3D	Integer	Number of additional diagnostic 3-d spectral output (0 = off; 1 = on)	0

Parameter	Type	Purpose	Default
NDL(NLEV)	Integer Array	Switch for diagnostic print out of a level (0 = off; 1 = on)	NLEV · 0
NHDIFF	Integer	Cut off wave number for horizontal diffusion	15
NHORDIF	Integer	Switch for horizontal diffusion (0 = off; 1 = on)	1
NTIME	Integer	Switch for CPU time diagnostics (0 = off; 1 = on)	0
NPERPETUAL	Integer	Switch for perpetual simulations (0 = annual cycle; >0 = day of the year)	0
DTEP	Real	Equator to pole temperature difference [K] for Newtonian cooling (usually not used)	0.0
DTNS	Real	North pole to south pole temperature difference [K] for Newtonian cooling (usually not used)	0.0
DTROP	Real	Tropopause height [m] for Newtonian cooling (usually not used)	12000.0
DTTRP	Real	Smoothing of the tropopause [K] for Newtonian cooling (usually not used)	2
TGR	Real	Surface temperature [K] for Newtonian cooling (usually not used)	280
TDISSD(NLEV)	Real Array	time scale [d] for the horizontal diffusion of divergence	NLEV · 0.2
TDISSZ(NLEV)	Real Array	time scale [d] for the horizontal diffusion of vorticity	NLEV · 1.1
TDISST(NLEV)	Real Array	time scale [d] for the horizontal diffusion of temperature	NLEV · 5.6
TDISSQ(NLEV)	Real Array	time scale [d] for the horizontal diffusion of moisture	NLEV · 5.6
PSURF	Real	Global mean sea level pressure [Pa]	101100.00
RESTIM(NLEV)	Real Array	Time scale [d] for Newtonian cooling (usually not used)	NLEV · 0.0

Parameter	Type	Purpose	Default
MARS	Integer	1: Set all parameters for Mars atmosphere	0
NGUI	Integer	Run with (1) or without (0) GUI	0
NOUTPUT	Integer	Global witch for enabling (1) or disabling (0) output to file puma_output	1
SELLON	Real	Longitude of soundings in the GUI	0.0
T0(NLEV)	Real Array	Reference temperature used in the discretization scheme	NLEV · 250.0
TFRC(NLEV)	Real Array	Time scale [d] for Rayleigh friction (0.0 = off)	NLEV · 0.0

Structure Internally, `plasim.f90` uses the FORTRAN-90 global common module `pumamod` from `plasimmod.f90`. After starting MPI, the main program `plasim` calls `prolog` for initializing the model. Then, `master` is called to do the time stepping. Finally, subroutine `epilog` finishes the run. In subroutine `prolog`, calls to different subroutines, which are part of `plasim.f90` or are provided by other modules, initialize various parts of the model: `gauaw` and `inilat` build the grid, `readnl` reads the namelist and sets some parameter according to the namelist input, `initpm` and `initsi` initialize some parameter for the physics and the semi implicit scheme, respectively. `outini` starts the output. If a file named **plasim_restart** exists all variables and arrays are read by `restart`, otherwise `initfd` sets the prognostic variables to their initial values. Calls to `miscini`, `fluxini`, `radini`, `rainini` and `surfini` start the initialization of the respective external modules. Finally, the global mean surface pressure is set according to PSURF (the observed value is 1011 hPa (Trenberth 1981) while 1013 is the ICAO standard) and the orography. Subroutine `master` controls the time stepping. First, if its not a restart, initial NKITS explicit forward timesteps are performed. The main loop is defined by calling `gridpointa` for the adiabatic tendencies, `spectrala` to add the adiabatic tendencies, `gridpointd` for the diabatic tendencies (which are computed by the external modules), `spectrald` to add the diabatic tendencies and the interface routines to the output module `outmod.f90`. The run is finalized by subroutine `epilog` which writes the restart records and calls the respective interface routines of the external modules.

2.10 plasimmod.f90

General The file `plasimmod.f90` contains the module `pumamod.f90` which declares all parameters and variables which may be used to share information between `plasim.f90` and other modules. No subroutines or programs are included.

Input/Output `pumamod.f90` does not use any extra input file or output file. No namelist input is required

Structure Internally, `plasimmod.f90` is a FORTRAN-90 module named *pumamod*. Names for global parameters, scalars and arrays are declared and, if possible, values are preset.

2.11 radmod.f90

General The module `radmod.f90` contains subroutines to compute radiative energy fluxes and the temperature tendencies due to long wave and short wave radiation. The interface to the main PLASIM module `plasim.f90` is given by the subroutines `radini`, `radstep` and `radstop` which are called in `plasim.f90` from the subroutines `prolog`, `gridpointd` and `epilog`, respectively.

Input/Output `radmod.f90` does not use an extra output file. If the Switch for ozone (NO₃, see namelist) is set to 2 (externally prescribed), the climatological cycle of the ozone distribution is read from the external file `surface.txt` which name is given in the namelist. The file format is formatted SERVICE format with (8I10) for the header and (8E12.6) for the fields. To assign the fields, the headers need to have the header information code 200, level going from 1 to NLEV and a date identifier of the form `yymmdd` or `mmdd` where `mm` goes from 01 to 12 (January to December). `radmod.f90` is controlled by the namelist `radpar` which is part of the namelist file `puma_namelist`:

Parameter	Type	Purpose	Default
NDCYCLE	Integer	Switch for diurnal cycle of insolation (0 = off, 1 = on)	0
NO3	Integer	Switch for ozone (0 = off, 1 = idealized distribution, 2 = externally prescribed)	1
CO2	Real	CO ₂ concentration [ppmv]	360.0
GSOL0	Real	Solar constant [W/m ²]	1367.0
IYRBP	Integer	Year PB (reference is 1950) to calculate orbit from	-50
NSWR	Integer	Switch for short wave radiation (0 = off, 1 = on)	1
NLWR	Integer	Switch for long wave radiation (0 = off, 1 = on)	1
NSOL	Integer	Switch for incoming solar radiation (0 = off, 1 = on)	1
NSWRCL	Integer	Switch for computed short wave cloud properties (0 = off, 1 = on)	1
NRSCAT	Integer	Switch for Rayleigh scattering (0 = off, 1 = on)	1
RCL1(3)	Real Array	Prescribed cloud albedos [frac.] for high, middle and low level clouds (spectral range 1)	0.15,0.30,0.60

Parameter	Type	Purpose	Default
RCL2(3)	Real Array	Prescribed cloud albedos [frac.] for high, middle and low level clouds (spectral range 2)	0.15,0.30,0.60
ACL2(3)	Real Array	Prescribed cloud absorptivities [frac.] for high, middle and low level clouds (spectral range 2)	0.05,0.10,0.20
CLGRAY	Real	Prescribed grayness of clouds (-1.0 = computed)	-1.0
TPOFMT	Real	Tuning for point of mean transmission	0.15
ACLLWR	Real	Mass absorption coefficient for clouds (long wave)	0.1
TSWR1	Real	Tuning of cloud albedo (spectral range 1)	0.035
TSWR2	Real	Tuning of cloud back scattering (spectral range 2)	0.04
TSWR3	Real	Tuning of cloud single scattering albedo (spectral range 2)	0.006
DAWN	Real	Threshold for zenith angle	0.0

Structure Internally, `radmod.f90` uses the FORTRAN-90 module `radmod`, which uses the global common module `pumamod` from `plasimmod.f90`. Additionally, the FORTRAN-90 module `orbparam` is used. Subroutine `radini` reads the namelist and, if the parallel version is used, distributes the namelist parameters to the different processes. Orbital parameters are computed by calling `orb_params`. If NO3 is set to 2, the ozone distribution is read from `surface.txt`. Subroutine `radstep` calls the subroutines `solang` and `mko3` to compute the cosine of the solar angle and the ozone distribution, respectively. The short wave radiative fluxes are calculate in `swr` while the long wave radiative fluxes are computed in `lwr`. Subroutine `radstop` is a dummy subroutine since there is nothing to do to finalize the computations in `radmod.f90`.

2.12 rainmod.f90

General The module `rainmod.f90` contains subroutines to compute large scale and convective precipitation and the related temperature tendencies. In addition, a parameterization of dry convective mixing of temperature and moisture is included and cloud cover is diagnosed. The interface to the main PLASIM module `plasim.f90` is given by the subroutines `rainini`, `rainstep` and `rainstop` which are called in `puma.f90` from the subroutines `prolog`, `gridpointd` and `epilog`, respectively.

Input/Output `rainmod.f90` does not use any extra input or output file and is controlled by the namelist `rainpar` which is part of the namelist file `puma_namelist`:

Parameter	Type	Purpose	Default
KBETTA	Integer	Switch for betta in Kuo parameterization (0 = off, 1 = on)	1
NPRL	Integer	Switch for large scale precipitation (0 = off, 1 = on)	1
NPRC	Integer	Switch for convective precipitation (0 = off, 1 = on)	1
NDCA	Integer	Switch for dry convective adjustment (0 = off, 1 = on)	1
NSHALLOW	Integer	Switch for shallow convection (0 = off, 1 = on)	1
RCRIT(NLEV)	Real Array	Critical relative humidity for cloud formation	computed
CLWCRIT1	Real	Critical vertical velocity for cloud formation [Pa/s] (not active if $CLWCRIT2 > CLWCRIT1$)	-0.1
CLWCRIT2	Real	Critical vertical velocity for cloud formation [Pa/s] (not active if $CLWCRIT2 > CLWCRIT1$)	0.0

Structure Internally, `rainmod.f90` uses the FORTRAN-90 module `rainmod`, which uses the global common module `pumamod` from `plasimmod.f90`. Subroutine `rainini` reads the namelist and, if the parallel version is used, distributes the namelist parameters to the different processes. Subroutine `rainstep` calls the subroutine `mkdqdtp` to obtain the adiabatic moisture tendencies in grid point space, which are needed for the Kuo parameterization. `kuo` is called to compute the convective precipitation and the respective tendencies. Dry convective adjustment is performed in `mkdca`. Large scale precipitation is computed in `mkdsp`. Finally, diagnostic clouds are calculated in `mkclouds`. Subroutine `radstop` is a dummy subroutine since there is nothing to do to finalize the computations in `radmod.f90`.

2.13 seamod.f90

General The module `seamod.f90` is the interface from the atmosphere to the ocean and the sea ice. The interface to the main PLASIM module `puma.f90` is given by the subroutines `seaini`, `seastep` and `seastop` which are called in `puma.f90` from the subroutines `prolog`, `gridpointd` and `epilog` respectively.

Input/Output `seamod.f90` reads different surface parameters either from the file `surface.txt` (see namelist) and the file `ocean_parameter` or from the restart file `sea_restart` which is written at the end of an integration.. The files formats are unformatted for the restart file, formatted SERVICE format with (8I10) for the header and (8E12.6) for the fields for `surface.txt` and formatted EXTRA format with (4I10) for the header and (6(1X,E12.6)) for the fields for `ocean_parameter`. The file `surface.txt` may include the following fields: The climatological annual cycle of the surface temperature [K] and the climatological annual cycle of the sea ice compactness [frac.]. To assign the fields, the headers need to have the header information code 169 for surface temperature and code 210 for the compactness (1 = ice; 0. = open water). a date identifier of the form `yymmdd` or `mddd` where `mm` goes from 1 to 12 (January to December) is required. Fields which are not needed will be skipped. The file `ocean_parameter` includes the following fields: The climatological annual cycle of the sea surface temperature [K], the climatological annual cycle of the mixed layer depth [m] and the climatological average of the deep ocean temperature [m]. To assign the fields, the order must be as described above (no header information is used). The restart file `sea_restart` contains all variables needed to continue the run. `seamod.f90` is controlled by the namelist `seapar` given in the namelist file `sea_namelist`:

Parameter	Type	Purpose	Default
ALBSEA	Real	Albedo for ice free ocean	0.069
ALBICE	Real	Maximum albedo for sea ice	0.7
DZ0SEA	Real	Minimum roughness length [m] for ice free ocean	$1.0 \cdot 10^{-5}$
DZ0ICE	Real	Roughness length [m] for sea ice	$1.0 \cdot 10^{-3}$
DRHSSEA	Real	Wetness factor for ice free ocean	1.0
DRHSICE	Real	Wetness factor for sea ice	1.0
NOCEAN	Integer	Switch for ocean model (0 = climatological SST, 1 = ocean model)	1
NICE	Integer	Switch for sea ice model (0 = climatological, 1 = sea ice model)	1

Parameter	Type	Purpose	Default
NCPL_ICE_OCEAN	Integer	ice-ocean coupling time steps	32
NCPL_ATMOS_ICE	Integer	ice atmosphere coupling time steps	1
TDEEPSEA	Real	Homogeneous deep ocean temperature [K]	0.0
DHICEMIN	Real	Minimum sea ice thickness [m]	0.1

Structure Internally, `seamod.f90` uses the FORTRAN-90 module `seamod`, which uses the global common module `pumamod` from `plasimmod.f90`. Subroutine `seaini` reads the namelist and, if the parallel version is used, distributes the namelist parameters to the different processes. If it is not a restart (i.e. if `NRESTART` from `inp` of `plasimmod.f90` is 0), the files `surface.txt` and `ocean_parameter` are being read. The climatological sea ice compactness is converted to a sea ice thickness as initial condition and additional surface parameters are set. If it is a restart, the restart file `sea_restart` is read. Subroutine `seastep` accumulates the variables used for the coupling between the atmosphere and the ocean. The coupling is done via the sea ice model. There is no direct connection between atmosphere and ocean model. If there is no sea ice, the coupling quantities are passed through the ice model without changes. Subroutine `seastop` finalizes the run and writes the restart records.

2.14 Sea ice and ocean modules

This section describes the modules that represent sea ice and ocean and the necessary interfaces between these modules and the atmospheric modules. Conceptually, the sea ice model lies inbetween the atmosphere model and the ocean model. Thus, the PUMA main part and the ocean model are both coupled to the sea ice model, but not directly to each other. The sea ice model decides whether a given gridpoint is covered with ice or not, in the latter case, it merely functions as passing the ocean fluxes to the atmosphere and vice versa. The parameters that are exchanged are listed in Table 2.1. The sea ice and ocean model use a time step of one day. Thus, atmospheric coupling to the sea ice model is performed every 32 time steps, while the sea ice and ocean model are coupled every time step. The coupling scheme is shown in Fig. 2.1. Fig. 2.2 shows how the subroutines are placed when no external coupler is used.

Parameter	Atmosphere \leftarrow \rightarrow Ice	Ice \leftarrow \rightarrow Ocean
Ice cover	\leftarrow	$-$
Ice thickness	\leftarrow	\rightarrow
Snow thickness	\leftarrow	\rightarrow
Surface temperature	\leftarrow	\leftarrow
Deep sea temperature	$-$	\leftarrow
Mixed layer depth	$-$	\leftarrow
Net precipitation, runoff	\rightarrow	\rightarrow
Salinity	$-$	\leftarrow
Melt and freeze volume	$-$	\rightarrow
Heat fluxes	\rightarrow	\rightarrow
d(Heat fluxes)/dT	\rightarrow	$-$
Radiation	\rightarrow	$-$
Wind stress	\rightarrow	\rightarrow

Table 2.1: Parameters to be exchanged between models. Arrows denote the direction in which the parameter is passed, e.g. the atmosphere receives ice cover information from the ice model.

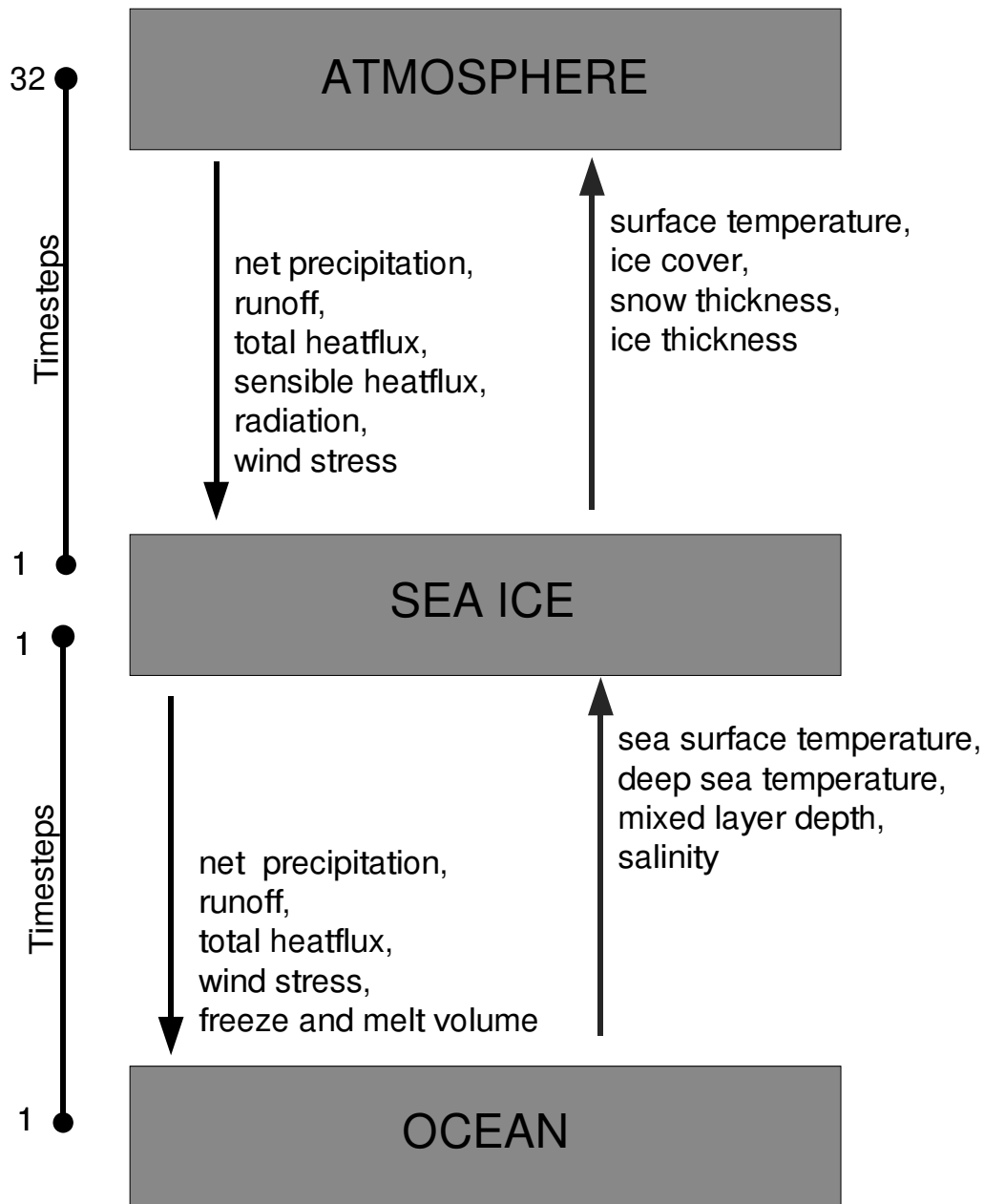


Figure 2.1: Schematic illustration of the model coupling.

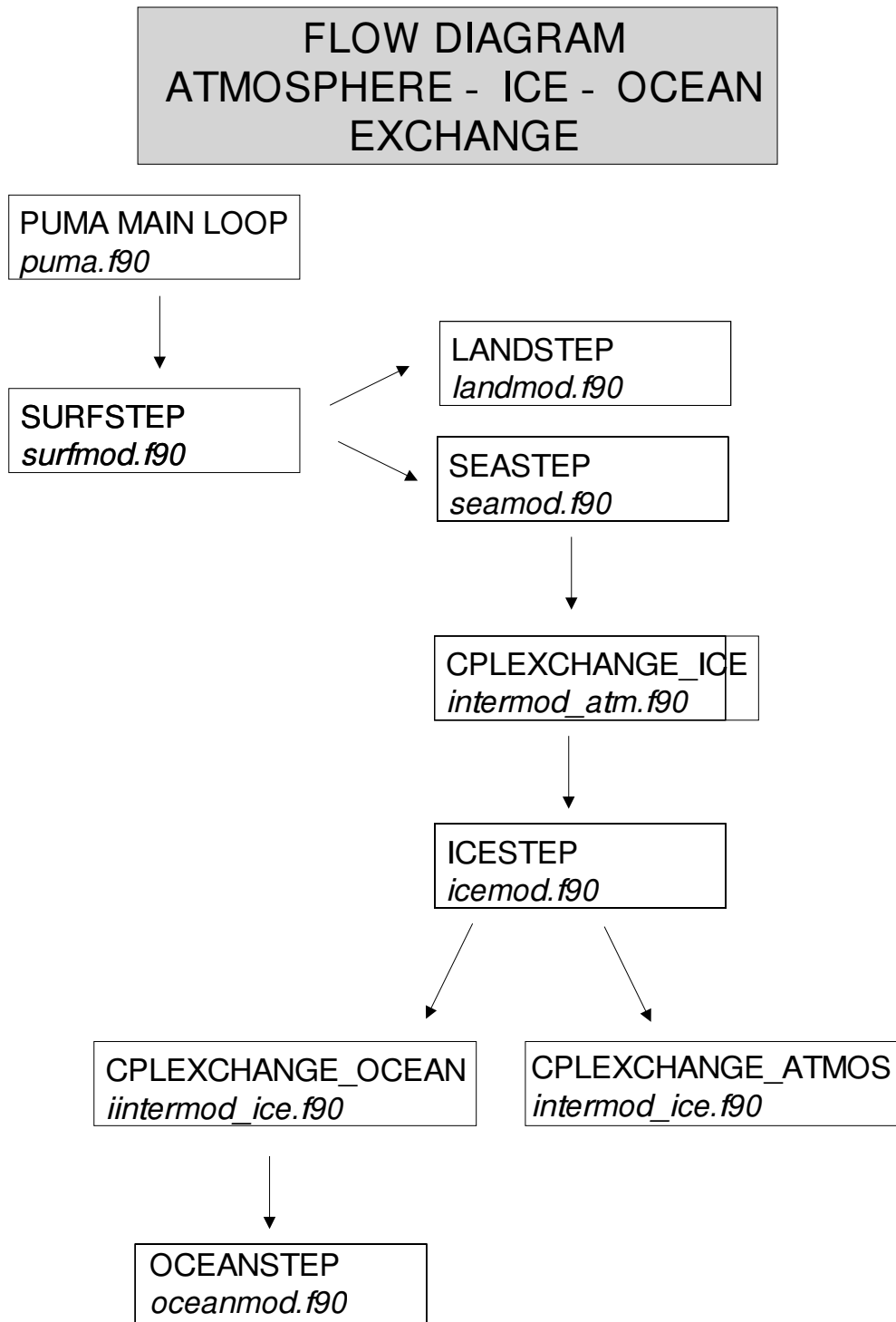


Figure 2.2: Subroutine flow when no external coupler is used.

2.15 icemod.f90

General The module `icemod.f90` contains subroutines to compute sea ice cover and thickness. The interface to the main PLASIM module is given by the subroutine `icestep`, which is called by `cplexchange_ice` (defined in `intermod_atm.f90`), which is called by `seastep` (defined in `seamod.f90`).

Input/Output `icemod.f90` requires the file `ice_flgcor` if `NFLXCORR` is set to a negative value. If `NOUTPUT` is set to 1, the output files **fort.75** containing global fields of ice model data and the file **fort.76** containing diagnostic ice data are produced (for details, see the reference manual). Both output files are in service format. The module is controlled by the namelist `icepar` in the file `ice_namelist`.

Parameter	Type	Purpose	default
NDIAG	INTEGER	Diagnostic output every NDIAG time steps	160
NOUT	INTEGER	Model data output every NOUT time steps	32
NOUTPUT	INTEGER	Icemodel output (0=no,1=yes)	1
NFLXCORR	INTEGER	Time constant for restoring (> 0), no flux correction (= 0), use flux-correction from file (< 0)	360 d

Structure `icemod.f90` uses the module `icemod` which is not dependent on the module `pumamod`. Subroutine `iceini` reads the namelist and, when required, the flux correction from the file `ice_flgcor`. Subroutine `icestep` calls `cplexchange_atmos` (defined in `intermod_ice`) to get the atmospheric forcing fields. If the `sea_namelist` parameter `NICE` is set to 1, the subroutine `subice` is called, which calculates ice cover and thickness. Otherwise, climatological data, interpolated to the current time step by `iceget` are used. If an ice cover is present, the surface temperature is calculated in `skintemp`. Otherwise, the surface temperature is set to the sea surface temperature calculated by the ocean model. Every `NCPL_ICE_OCEAN` (defined in `sea_namelist`) time steps, the external subroutine `cplexchange_ocean` (defined in `intermod_ice`) is called to pass the atmospheric forcing to and retrieve oceanic data from the ocean module `oceanmod.f90`. The oceanic data is used for ice calculations in the next time step.

2.16 oceanmod.f90

General The module `oceanmod.f90` contains a mixed layer ocean model, i.e. subroutines to compute sea surface temperature and mixed layer depth. The interface to the main PLASIM module is via the module `icemod.f90` given by the subroutine `oceanstep`, which is called by `cplexchange_ocean` (defined in `intermod_ice`).

Input/Output `oceanmod.f90` requires the file `ocean_fluxcor` if `NFLXCORRSST` or `NFLXCORRMLD` is set to a negative value. If `NOUTPUT` is set to 1, the output file `fort.31` containing global fields of ocean model data in service format is produced (for details, see the ice modul section of the reference guide). The module is controlled by the namelist `oceanpar` in the file `ocean_namelist`.

Parameter	Type	Purpose	default
NDIAG	INTEGER	Diagnostic output every NDIAG time steps	480
NOUT	INTEGER	Model data output every NOUT time steps	32
NOUTPUT	INTEGER	Oceanmodel output (0=no,1=yes)	1
NFLXCORRMLD	INTEGER	Time constant for restoring mixed layer depth (> 0), no flux correction (= 0), use fluxcorrection from file (< 0)	60 d
NFLXCORRSST	INTEGER	Time constant for restoring sea surface temperature (> 0), no flux correction (= 0), use fluxcorrection from file (< 0)	60 d

Structure `oceanmod.f90` uses the module `oceanmod` which is not dependent on the module `pumamod`. Subroutine `oceanini` reads the namelist and, when required, the flux corrections from the file `ocean_fluxcor`. Subroutine `oceanstep` calls `mixocean`, which calculates mixed layer depth and temperature. If an ice cover is present, mixed layer depth is set to the climatological value and the sea surface temperature is set to the freezing temperature. For details of the mixed layer model, see the Planet Simulator Reference Manual.

Chapter 3

Parallel Program Execution

3.1 Concept

The **Planet Simulator** is coded for parallel execution on computers with multiple CPU's or networked machines. The implementation uses MPI (Message Passage Interface), that is available for nearly every operating system <http://www.mcs.anl.gov/mpi>.

In order to avoid maintaining two sets of source code for the parallel and the single CPU version, all calls to the MPI routines are encapsulated into a module. Users, that want to compile and execute the parallel version use the module **mpimod.f90** and the commands **mpif90** for compiling and **mpirun** for running.

If MPI is not implemented or the single CPU version is sufficient, **mpimod_stub.f90** is used instead of **mpimod.f90**. Also remove or comment the line:

```
!      use mpi
```

and set the number of processors to 1:

```
parameter(NPRO = 1)
```

3.2 Parallelization in Gridpoint Domain

The data arrays in gridpoint domain are either three-dimensional e.g. `gt(NLON, NLAT, NLEV)` referring to an array organized after longitudes, latitudes and levels, or two-dimensional, e.g. `gp(NLON, NLAT)`. The code is organized such, that there are no dependencies in latitudinal direction, while in gridpoint domain. Such dependencies are resolved during the Legendre-Transformations. So the the partitioning of the data is done in latitudes. The program can use as many CPU's as latitudes with the extreme of every CPU doing the computations for a single latitude. There is the restriction however, that the number of latitudes (`NLAT`) divided by the number of processors (`NPRO`), giving the number of latitudes per process (`NLPP`) must have zero remainder. E.g. A T31 resolution uses $NLAT = 48$. Possible values for `NPRO` are then 1, 2, 3, 4, 6, 8, 12, 16, 24, and 48.

All loops dealing with a latitudinal index look like:

```
do jlat = 1 , NLPP
    ....
enddo
```

There are, however, many subroutines, with the most prominent called **calcgp**, that can fuse latitudinal and longitudinal indices. In all these cases the dimension `NHOR` is used. `NHOR` is defined as: $NHOR = NLON * NLPP$ in the `pumamod` - module. The typical gridpoint loop that looks like:

```

do jlat = 1 , NLPP
  do jlon = 1 , NLON
    gp(jlon,jlat) = ...
  enddo
enddo

```

is then replaced by the faster executing loop:

```

do jhor = 1 , NHOR
  gp(jhor) = ...
enddo

```

3.3 Parallelization in Spectral Domain

The number of coefficients in spectral domain (NRSP) is divided by the number of processes (NPRO) giving the number of coefficients per process (NSPP). The number is rounded up to the next integer and the last process may get some additional dummy elements, if there is a remainder in the division operation.

All loops in spectral domain are organized like:

```

do jsp = 1 , NSPP
  sp(jsp) = ...
enddo

```

3.4 Synchronization points

All processes must communicate and have therefore to be synchronized at following events:

- Legendre-Transformation: This involves changing from latitudinal partitioning to spectral partitioning and such some gather and scatter operations.
- Inverse Legendre-Transformation: The partitioning changes from spectral to latitudinal by using gather, broadcast, and scatter operations.
- Input-Output: All read and write operations must be done only by the root process, who gathers and broadcasts or scatters the information as desired. Code that is to be executed by the root process exclusively is written like:

```

if (mypid == NROOT) then
  ...
endif

```

NROOT is typically 0 in MPI implementations, mypid (My process identification) is assigned by MPI.

3.5 Source code

It needs some discipline in order to maintain parallel code. Here are the most important rules for changing or adding code to the **Planet Simulator**:

- Adding namelist parameters: All namelist parameters must be broadcasted after reading the namelist. (Subroutines mpbci, mpbcr, mpbcin, mpbcn)
- Adding scalar variables and arrays: Global variables must be defined in a module header and initialized.
- Initialization code: Initialization code, that contains dependencies on latitude or spectral modes must be done by the root process only and then scattered from there to all child processes.
- Array dimensions and loop limits: Always use parameter constants (NHOR, NLAT, NLEV, etc.) as defined in pumamod.f90 for array dimensions and loop limits.
- Testing: After significant code changes the program should be tested in single and in multi-CPU configuration. The results of a single CPU run is usually not exactly the same as the result of a multi-CPU run due to effects in rounding. But the results should show only small differences during the first timesteps.
- Synchronization points: The code is optimized for parallel execution and minimizes therefore communication overhead. The necessary communication code is grouped around the Legendre-transformations. If more scatter/gather operations or other communication routines are to be added, they should be placed just before or after the execution of the calls to the Legendre-Transformation. Any other place would degrade the overall performance in introducing additional process synchronization.

Chapter 4

Graphical User Interface

4.1 Graphical user interface (GUI)

The **Planet Simulator** may be used in the traditional fashion, with shell scripts, batch jobs, and network queuing systems. This is acceptable for long running simulations on complex machines and number-crunchers, like vector- computers, massive-parallel-computers and workstation clusters. There is now, however, a much more convenient method by using a graphical user interface (GUI) for model setup with parameter configurations and for interaction between user and model.

The **Planet Simulator** is configured and setup by the first GUI module named MoSt (Model Starter, screenshot in 4.1). MoSt is the fastest way to get the model running. It gives access to the most important parameters of the model preset to the most frequently used values. The model can be started with a mouse click on the button labelled "Save & Run" either with the standard parameter setting or after editing some of the parameters in the MoSt window. Some parameters, like horizontal and vertical resolution, or the number of processors, require the building (compile, link and load) of new executables. MoSt achieves this by generating and executing build scripts, that perform the necessary code changes and create the required executable. Other parameters define startup- and boundary conditions or settings for parameterisations. They can be edited in MoSt and, after a check for correct range and consistency with other parameters, are written to the model's namelist file.

Depending on all settings MoSt generates a runsript for the simulation. The user has the choice of leaving MoSt and continue with the simulation under control of a GUI right away, or to exit MoSt with the scripts prepared to run. The second alternative is useful for users, who want to modify the setup beyond the scope of MoSt or want to run the Planet Simulator without GUI.

There's also a simple graphical editor for topography. Check the box Orography and then use the mouse to mark rectangular areas in the topography display. Enter a value for rising (positive) or lowering the area and press the button labelled **Preprocess**. The preprocessor will be built and executed, a new topography will be computed and written to a start file.

Another editor is the mode editor for spherical harmonics. Green modes are enabled, red modes are disabled. This feature can be used to make runs with only certain modes of spherical harmonics being active. MB1, MB2, MB3 refer to the left, middle, and right mouse button. You may toggle individual modes or whole lines and columns. Currently this mode editor can only be used for **Planet Simulator** in the T21 resolution.

The GUI for running the **Planet Simulator** (screenshot in 4.2) has two main purposes. The first one is to display model arrays in suitable representations. Current implementations are:

- Zonal mean cross sections

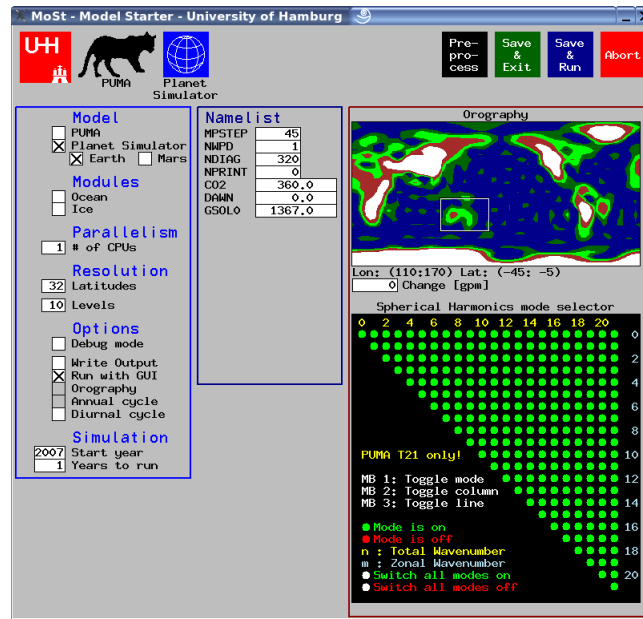


Figure 4.1: Screenshot of Model Starter (MoSt)

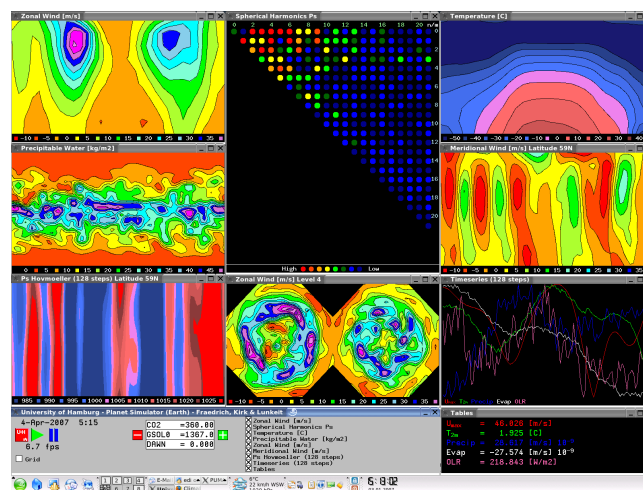


Figure 4.2: Screenshot of Graphical User Interface (GUI)

- Horizontal global fields in cylinder projection
- Horizontal global fields in polar projection
- Time-longitude (Hovmoeller) diagrams
- Amplitudes of coefficients of spherical harmonics
- Time series
- Numerical values

In case of horizontal global grids pressing the MMB (Middle Mouse Button) toggles between cylinder and polar projection. If the grid is just one level from many of a three dimensional field like u or v , the level shown can be decreased by the LMB or increased by the RMB. For Hovmoeller and longitude height sections the LMB and RMB can be used to select the latitude.

The second purpose is the interaction part of the GUI, which allows the user to change selected model variables during the model run. It is not necessary, though possible, to pause the model while changing variables. Changes to model variables are, of course, monitored in the outputfile and checked by GUI for the appropriate range of values and maximum possible change per timestep because, otherwise, a rapid parameter change or a choice of values beyond the normal range may blow up the model.

All model variables, which are candidates for the display or interactive changes, have a special code to communicate with the Planet Simulator. The experienced modeller can add new code for more variables using the existing communication code as template. Thus all model fields or even fields received via coupling with other models can be put on the GUI display.

Both, MoSt and GUI are implemented using the Xlib (X11R5), which is a library of routines for graphics and event communication. As this library is part of every UNIX/Linux operating system and base of all desktop environments, there is no need to install additional software for running MoSt and GUI. Another important property of Xlib is the full network transparency. The display of MoSt and GUI is not locked to the machine running the programs or the model. In fact, the best performance is obtained in running the Planet Simulator on two or four CPUs of a remote server while displaying the GUI on the user's workstation. In summarizing, the MoSt and GUI programs automate many tedious tasks, minimize the time to become familiar with the Planet Simulator, and make debugging and parameter tuning much easier. More kinds of presentations, coordinate projections and interactivity are being developed. A graphical preprocessor with editor for boundary conditions and a graphical postprocessor are future expansions to build an almost complete environment for modellers.

4.2 GUI configuration

On initialization the GUI reads its configuration from a file **GUI.cfg** which must be present in the current directory. MoSt copies the file **GUI.cfg** from the `../dat/` directory to the run directory while building the **Planet Simulator**. After reading **GUI.cfg** an attempt is made to read the file **GUI_last_used.cfg**. This file is always written at the end of a GUI controlled simulation. So one may rearrange and position GUI windows during a run and the new layout will be saved to the file **GUI_last_used.cfg**. In order to make this user layout default for following runs, just copy this file like:

```
Most15/plasim/run$ cp ../dat/GUI.cfg ../dat/GUI.cfg.old
Most15/plasim/run$ cp GUI_last_used.cfg ../dat/GUI.cfg
```

MoSt will then copy your new layout to the run directory at the next invocation.

The **GUI.cfg** is a text file that may be also edited manually. There is a section for each window (counting from 0 to 8) which looks like:

```
[Window 00]                <- window number (0..8)
Array:CSU                  <- array name
Plot:ISOCS                 <- picture type
Palette:U                  <- colour palette
Title:Zonal Wind [m/s]    <- window title
Geometry: 529 299 2 3     <- width height left top

[Window 01]
Array:SPAN
Plot:ISOSH
Palette:AMPLI
Title:Spherical Harmonics Ps
Geometry: 529 299 535 3

...
```

Possible values for these items are:

4.2.1 Array

Name	Description
CSU	Cross Section U - Zonal mean zonal wind
CSV	Cross Section V - Zonal mean meridional wind
CST	Cross Section T - Zonal mean temperature
SPAN	Spherical harmonics coefficients of surface pressure
GU	Three dimensional grid of zonal wind
GV	Three dimensional grid of meridional wind
GP	Grid of surface pressure
DQVI	Vertically integrated humidity
GUCOL	sounding of eastward wind at a grid point
GVCOL	sounding of northward wind a grid point
GTCOL	sounding of temperature at a grid point
DQCOL	sounding of humidity at a grid point
DQLCOL	sounding of liquid water at a grid point
DCCCOL	sounding of cloud cover at a grid point
SCALAR	Selected scalars for timeseries and tables

4.2.2 Plot

Name	Description
ISOHOR	Isolines and colouring of horizontal grids
ISOCS	Isolines and colouring of cross sections
ISOHOV	Colouring of Hovmoeller diagram
ISOTS	Timeseries
ISOTAB	Tables
ISOSH	Coloured amplitudes
ISOLON	Isolines and colouring of longitude height section
ISOCOL	vertical Hovmoeller diagram for soundings

4.2.3 Palette

Name	Range	Description
AUTO	automatic	rainbow colours
U	-10 .. 50	rainbow colours
V	-10 .. 10	rainbow colours
T	-50 .. 50	blue - red
P	985 .. 1025	blue - red
Q	0 .. 60	rainbow colours
DCC	0 ..100	rainbow colours
MARST	-90 .. 0	blue -red
AMPLI	0 .. 12	blue - green -red
VEG	0 .. 100	shades of green

4.2.4 Title

The title item may contain any text, but keep it short, the length of the window's title bar is limited. The words *Latitude* and *Level* have special features in conjunction with threedimensional arrays, where the user may scroll the level or latitude. The GUI will insert the level number after the word *Level* or the latitude after the word *Latitude*.

4.2.5 Geometry

The four integers following the geometry item describe the size and screen position of the window. The first two parameters refer to width and height in screen pixel. These are the sizes of the inner window, title bar, border and other decorations are not counted. The third and fourth parameter set the coordinates of the upper left corner of the window x and y, again without borders. If the geometry item is not defined, the GUI will initialize the window's geometry depending on the screen size.

Chapter 5

Postprocessor Pumaburner

5.1 Introduction

The **Pumaburner** is a postprocessor for the **Planet Simulator** and the **PUMA** model family. It's the only interface between *raw* model data output and diagnostics, graphics, and user software.

The output data of the **Planet Simulator** are stored as packed binary (16 bit) values using the model representation. Prognostic variables like temperature, divergence, vorticity, pressure, and humidity are stored as coefficients of spherical harmonics on σ levels. Variables like radiation, precipitation, evaporation, clouds, and other fields of the parameterization package are stored on Gaussian grids.

The tasks of the **Pumaburner** are:

- Unpack the *raw* data to full real representation.
- Transform variables from the model's representation to a user selectable format, e.g. grids, zonal mean cross sections, fourier coefficients.
- Calculate diagnostic variables, like vertical velocity, geopotential height, wind components, etc.
- Transform variables from σ levels to user selectable pressure levels.
- Compute monthly means and standard deviations.
- Write selected data either in SERVICE, GRIB, or NetCDF format for further processing.

5.2 Usage

```
pumaburn4 [options] InputFile OutputFile <namelist >printout
  option -h : help (this output)
  option -c : print available codes and names
  option -d : debug mode (verbose output)
  option -g : Grib   output (override namelist option)
  option -n : NetCDF output (override namelist option)
  option -m : Mean=1 output (override namelist option)
  InputFile : Planet Simulator or PUMA data file
  OutputFile : GRIB, SERVICE, or NetCDF format file
  namelist  : redirected <stdin>
  printout  : redirected <stdout>
```

5.3 Namelist

The namelist values control the selection, coordinate system and output format of the post-processed variables. Names and values are not case sensitive. You can assign values to the following names:

Name	Def.	Type	Description	Example
HTYPE	S	char	Horizontal type	HTYPE=G
VTYPE	S	char	Vertical type	VTYPE=P
MODLEV	0	int	Model levels	MODLEV=2,3,4
hPa	0	real	Pressure levels	hPa=500,1000
CODE	0	int	ECMWF field code	CODE=130,152
GRIB	0	int	GRIB output selector	GRIB=1
NETCDF	0	int	NetCDF output selector	NETCDF=1
MEAN	1	int	Compute monthly means	MEAN=0
HHMM	1	int	Time format in Service format	HHMM=0
HEAD7	0	int	User parameter	HEAD7=0815
MARS	0	int	Use constants for planet Mars	MARS=1
MULTI	0	int	Process multiple input files	MULTI=12

5.4 HTYPE

HTYPE accepts the first character of the following string. Following settings are equivalent: HTYPE = S, HTYPE=Spherical Harmonics HTYPE = Something. Blanks and the equal-sign are optional.

Possible Values are:

Setting	Description	Dimension for T21 resolution
HTYPE = S	Spherical Harmonics	(506):(22 * 23 coefficients)
HTYPE = F	Fourier Coefficients	(32,42):(latitudes,wavenumber)
HTYPE = Z	Zonal Means	(32,levels):(latitudes,levels)
HTYPE = G	Gauss Grid	(64,32):(longitudes,latitudes)

5.5 VTYPE

VTYPE accepts the first character of the following string. Following settings are equivalent: VTYPE = S, VTYPE=Sigma VTYPE = Super. Blanks and the equal-sign are optional.

Possible Values are:

Setting	Description	Remark
VTYPE = S	Sigma (model) levels	Some derived variables are not available
VTYPE = P	Pressure levels	Interpolation to pressure levels

5.6 MODLEV

MODLEV is used in combination with **VTYPE = S**. If **VTYPE** is not set to Sigma, the contents of **MODLEV** are ignored. **MODLEV** is an integer array that can get as many values as there are levels in the model output. The levels are numbered from top of the atmosphere to the bottom. The number of levels and the corresponding sigma values are listed in the pumaburner printout. The outputfile orders the level according to the **MODLEV** values. **MODLEV=1,2,3,4,5** produces an output file of five model levels sorted from top to bottom, while **MODLEV=5,4,3,2,1** sorts them from bottom to top.

5.7 hPa

hPa is used in combination with **VTYPE = P**. If **VTYPE** is not set to Pressure, the contents of **hPa** are ignored. **hPa** is a real array that accepts pressure values with the units hectoPascal or millibar. All output variables will be interpolated to the selected pressure levels. There is no extrapolation on the top of the atmosphere. For pressure values, that are lower than that of the model's top level, the top level value of the variable is taken. The variables temperature and geopotential height are extrapolated if the selected pressure is higher than the surface pressure. All other variables are set to the value of the lowest mode level for this case. The outputfile contains the levels in the same order as set in **hPa**. Example: `hpa = 100,300,500,700,850,900,1000`.

5.8 MEAN

MEAN can be used to compute montly means and/or deviations. The **Pumaburner** reads date and time information from the model file and handles different lengths of months and output intervals correctly.

Setting	Description
MEAN = 0	Do no averaging - all terms are processed.
MEAN = 1	Compute and write monthly mean fields. Not for spherical harmonics, Fourier coefficients or zonal means on sigma levels.
MEAN = 2	Compute and write monthly deviations. Not for spherical harmonics, Fourier coefficients or zonal means on sigma levels. Deviations are not available for NetCDF output.
MEAN = 3	Combination of MEAN=1 and MEAN=2. Each mean field is followed by a deviation field with an identical header record. Not for spherical harmonics, Fourier coefficients or zonal means on sigma levels.

5.9 Format of output data

The **pumaburner** supports three different output formats:

- **GRIB** (GRIdded Binary) WMO standard for gridded data.
- **NetCDF** (Network Common Data Format)
- **Service** Format for user readable data (see below).

For more detailed descriptions see for example:

<http://www.nws.noaa.gov/om/ord/iob/NOAAPORT/resources/>

Setting	Description	
GRIB = 1	NetCDF = 0	The output file is written GRIB format. This option can be used only for HTYPE=Spherical Harmonics or HTYPE=Gauss Grid.
GRIB = 0	NetCDF = 1	The output file is written in NetCDF format. This option can be used for HTYPE=Gauss Grid only.
GRIB = 0	NetCDF = 0	The output file is written in Service format. This is the preferred format for user programs. For a detailed description see the following section.
GRIB = 1	NetCDF = 1	Illegal combination.

5.10 SERVICE format

The SERVICE format uses the following structure: The whole file consists of pairs of header records and data records. The header record is an integer array of 8 elements.

```

head(1) = ECMWF field code
head(2) = modellevel or pressure in [Pa]
head(3) = date [yyymmdd] (yyymm00 for monthly means)
head(4) = time [hhmm] or [hh] for HHMM=0
head(5) = 1. dimension of data array
head(6) = 2. dimension of data array
head(7) = may be set with the parameter HEAD7
head(8) = experiment number (extracted from filename)

```

Example for reading the SERVICE format (GRIB=0 , NETCDF=0)

```

INTEGER HEAD(8)
REAL    FIELD(64,32)    ! dimensions for T21 grids
READ (10,ERR=888,END=999) HEAD
READ (10,ERR=888,END=999) FIELD
....
888 STOP 'I/O ERR'
999 STOP 'EOF'
....

```

5.11 HHMM

Setting	Description
HHMM = 0	head(4) shows the time in hours (HH).
HHMM = 1	head(4) shows the time in hours and minutes (HHMM).

5.12 HEAD7

The 7th. element of the header is reserved for the user. It may be used for experiment numbers, flags or anything else. Setting HEAD7 to a number exports this number to every header record in the output file (SERVICE format only).

5.13 MARS

This parameter is used for processing simulations of the Mars atmosphere. Setting MARS=1 switches gravity, gas constant and planet radius to the correct values for the planet Mars.

5.14 MULTI

The parameter MULTI can be used to process a series of input data within one run of the pumaburner. Setting MULTI to a number (n) tells the pumaburner to process (n) input files. The input files must follow one of the following two rules:

- YYMM rule: The last four characters of the filename contain the data in the form YYMM.

- .NNN rule: The last four characters of the filename consist of a dot followed by a 3-digit sequence number.

Examples:

Namelist contains MULTI=3

```
Command: pumaburn <namelist >printout run.005 out
pumaburn processes the files <run.005> <run.006> <run.007>
```

Namelist contains MULTI=4

```
Command: pumaburn <namelist >printout exp0211 out
pumaburn processes the files <exp0211> <exp0212> <exp0301> <exp0302>
```

5.15 Namelist example

```
VTYPE = Pressure
HTYPE = Grid
CODE = 130,131,132
hPa = 200,500,700,850,1000
MEAN = 0
GRIB = 0
NETCDF = 0
```

This namelist will write Temperature(130), u(130) and v(131) on pressure levels 200hPa, 500hPa, 700hPa, 850hPa and 1000hPa. The output interval is the same as found on the model data, e.g. every 12 or every 6 hours (MEAN=0). The output format is SERVICE format.

5.16 Troubleshooting

If the pumaburner reports an error or doesn't produce the expected results, try the following:

- Check your namelist, especially for invalid codes, types and levels.
- Run the pumaburner in debug-mode by using the option -d. Example:

```
pumaburn <namelist >printout -d data.in data.out
```

This will print out some details like parameters and memory allocation during the run. The additional information may help to detect the problem.

- Not all combinations of HTYPE, VTYPE, and CODE are valid. Try to use HTYPE=Grid and VTYPE=Pressure before switching to exotic parameter combinations.

Chapter 6

Graphics

6.1 Grads

In this section, visualisation using the graphics package GrADS is described. A useful Internet site for reference and installation instructions is

<http://grads.iges.org/grads/grads.html>.

Latest versions of GrADS can handle data in NETCDF format (via the command `sdfopen`), GRIB, HDF-SDS, and in its native binary format. The native format can conveniently be derived from SERVICE format. In the following it is assumed that the PUMA output has been converted to SERVICE format with the `pumaburner` and the resulting file is called `puma.srv`. Monthly mean data is either obtained directly from the `pumaburner` (`namelist` parameter `MEAN=1`, see section 5) or via a PINGO command:

```
srv monmeans puma.srv puma_m.srv
```

Information on the PINGO package can be found in DKRZ report 11 at

<http://www.mad.zmaw.de/Pingo/repdl.html>.

The SERVICE file has to be converted to GrADS's native format by the command:

```
srv2gra puma_m.srv
```

which results in the files `puma_m.gra` and `puma_mctl`. The first file contains the data, the latter one information on the grid, time steps, and variable names. The program `srv2gra` is one of the postprocessing tools available at

<http://puma.dkrz.de/puma/download/map/>.

If you chose to compile it yourself, please read the comments in the first few lines of the program text. Sometimes the `srv2gra` tool has difficulties to calculate an appropriate time increment from the date headers of the data records, so you should check this. In this example the file `puma_mctl` should look like this:

```
DSET ^puma_m.gra
UNDEF 9e+09
XDEF      64 LINEAR   0.0000   5.6250
OPTIONS YREV
YDEF      32 LEVELS
  -85.7606  -80.2688  -74.7445  -69.2130  -63.6786  -58.1430  -52.6065  -47.0696
```

```

-41.5325  -35.9951  -30.4576  -24.9199  -19.3822  -13.8445  -8.3067  -2.7689
   2.7689   8.3067  13.8445  19.3822  24.9199  30.4576  35.9951  41.5325
  47.0696  52.6065  58.1430  63.6786  69.2130  74.7445  80.2688  85.7606
ZDEF  1 LINEAR 1 1
TDEF 12 LINEAR 00:00Z01jan0001          1mo
VARS  3
c139   0 99   139   0   0
c151   0 99   151   0   0
c175   0 99   175   0   0
ENDVARS

```

Here, the line starting with `TDEF` ends with `1mo`, since we are handling monthly mean data. When the PUMA output is used without averaging, this should correspond to the output interval given by the `nafter` variable used in the `namelist` of your PUMA run (see section C). The number of variables depends on how the `pumaburner` was called. In this example, only 3 variables were processed, i.e. the surface temperature (`c139`), the sea level pressure (`c151`) and the albedo (`c175`; refer to appendix B for a list of codes).

The GrADS program is started by typing `grads` in a terminal window. Then, data is visualised either by typing commands line-by-line, or, preferably, by using scripts. The following script, called `tglob.gs`, displays the monthly mean surface temperature:

```

# tglob.gs
function pass(m)
'reinit'
'open puma_m'
'enable print print.mf'
'set t 'm
'c'
'set gxout shaded'
'd (c139-273.16)'
'cbar.gs'
'set gxout contour'
'd (c139-273.16)'
'draw title Surface Temperature (deg C) month 'm
'print'
'disable print'
'!gxps -i print.mf -o tglob'm'.ps'

```

The variable `m` at the beginning of the script defines the month which should be displayed. It is passed from the terminal with the script call. Note that in this line, no quotation marks are present, since only GrADS specific commands are framed by quotation marks. Script commands, like variable definitions, if-clauses etc. are used without quotation marks. The script is executed by typing its name without the ending and the number of the month to be shown. For example, `tglob 7` displays the monthly mean surface temperature in July. The resulting output file is called `tglob7.ps`.

The following script `thh` displays the time dependent surface temperature of Hamburg. Here, two variables are passed to GrADS, the first and last day to plot (note that here, the file `puma.gra` is opened, which contains data on a daily basis). The call `thh 91 180` displays the surface temperature of Hamburg for the spring season from April 1st to June 30th.

```

# thh.gs

```

```
function pass(d1 d2)
'reinit'
'open puma'
'enable print print.mf'
'set lat 53'
'set lon 10'
'set t 'd1' 'd2'
'c'
'd (c139-273.16)''
'draw title Surface Temperature (deg C) in Hamburg'
'print'
'disable print'
'!gxps -i print.mf -o thh.ps'
```

It is possible to have more than one figure in a plot, which is illustrated in the following script. It plots seasonal means of the sea level pressure. The data file is prepared like this:

```
srv selcode,151 puma.srv slp.srv
srv seamean slp.srv slp_sm.srv
srv2gra slp_sm.srv
```

The commands `set vpage` sets virtual pages inside the graphic window. The full window is 11 inch wide and 8.5 inch high, so `set vpage 0 5.5 4.25 8.5` defines the upper left corner. If `setlevs=1` is specified, the pressure levels as given are used. Otherwise, GrADS defines contour levels depending on the data set.

```
# slp_sm.gs
setlevs=1
'reinit'
'open slp_sm'
'enable print print.mf'
'c'
'set vpage 0 5.5 4.25 8.5'
'set gxout contour'
if (setlevs=1)
'set clevs 990 995 1000 1005 1010 1015 1020'
endif
'set ccols 1'
'set grads off'
'set t 1'
'd c151/100'
'draw title SLP [hPa] yr 'ny' DJF'
'set vpage 5.5 11 4.25 8.5'
'set gxout contour'
if (setlevs=1)
'set clevs 990 995 1000 1005 1010 1015 1020'
endif
'set ccols 1'
'set grads off'
'set t 2'
```

```
'd c151/100'
'draw title yr 'ny' MAM'
'set vpage 0 5.5 0 4.25'
'set gxout contour'
if (setlevs=1)
'set clevs 990 995 1000 1005 1010 1015 1020'
endif
'set ccols 1'
'set grads off'
'set t 3'
'd c151/100'
'draw title yr 'ny' JJA'
'set vpage 5.5 11 0 4.25'
'set gxout contour'
if (setlevs=1)
'set clevs 990 995 1000 1005 1010 1015 1020'
endif
'set ccols 1'
'set grads off'
'set t 4'
'd c151/100'
'draw title yr 'ny' SON'
'print'
'disable print'
'!gxps -c -i print.mf -o slp_sm.ps'
```

6.2 Vis5D

“Vis5D is a system for interactive visualization of large 5-D gridded data sets such as those produced by numerical weather models. One can make isosurfaces, contour line slices, colored slices, volume renderings, etc of data in a 3-D grid, then rotate and animate the images in real time. There’s also a feature for wind trajectory tracing, a way to make text annotations for publications, support for interactive data analysis, etc.”

from the Vis5D home page,
<http://www.ssec.wisc.edu/~billh/vis5d.html>

This powerful visualisation tool together with its documentation is available through the above home page. Vis5D uses its own data format which makes it necessary to transform your data. Depending on their format and the flowchart on <http://puma.dkrz.de/puma/download/map/> you have the following choices: If

- *your data is raw PUMA output,*
you need to process it with the `pumaburner` postprocessor (see section 5) in order to transform it to either NETCDF (option `-n` or namelist parameter `NETCDF=1`) or GRIB (option `-g` or namelist parameter `GRIB=1`) and proceed from there.
- *your data is in SERVICE format,*
you need to convert it to either GRIB, for instance with the PINGOs:

```
grb copy2 data.srv data_with_grib_metainfo.grb output.grb,
```

or NETCDF, using the program `puma2cdf`, which is available with the PUMA postprocessing tools. Despite of its name this program cannot process raw PUMA output but takes SERVICE format as input. It can as well be called as `srv2cdf` which changes its behaviour: oddities of model output such as the existence of February, 30th are then no longer removed. Once the format is changed proceed from there.

- *your data is in NETCDF format,*
it can easily transformed to Vis5D's native format by means of the program `cdf2v2d`, which is available with the PUMA postprocessing tools.
- *your data is in GRIB format,*
you can find a transformation tool named `Grib2V5d` at <http://grib2v5d.sourceforge.net> which offers various practical features.

Once the conversion to Vis5D's native format is achieved please follow the instructions from the Vis5D documentation or, if Vis5D is already installed on your system, try finding your own way by typing:

```
vis5d my_data.v5d
```


Chapter 7

Column Mode and Soundings

7.1 Setup

The column mode of the the Planet Simulator is an integral part of the full Planet Simulator and not a stand alone model. The advantage of this approach is that all options and modules available in the full model are automatically included in the column mode and that no extra maintenance is necessary.

Technically this is realized by switching off horizontal advection and diffusion which leaves us with an independent column for each grid point. The inclusion of additional options for boundary conditions allows for runs with synchronized columns.

Running the Model at the lowest resolution (T1) with 8 synchronized columns is efficient enough to run very fast column integrations. Using a standard setup with 10 atmospheric layers and a mixed layer ocean one can simulate more than 33500 years per day on a single processor PC (3.3 GHZ CPU).

To make full use of the computer power one can setup an ensemble of columns by specifying boundary conditions for every grid point separately.

7.1.1 Basic switches for column setup

We introduce the macro switch **COLUMN** in the namelist *inp* which is part of the namelist **puma_namelist**. By setting **COLUMN = 1** a default column mode is initialized by setting **YMODE = "Column"**, **KICK = 0**, **NADV = 0** and **NHORDIF = 0**. One can customize the column setup by keeping the default value **COLUMN = 0** and by setting the other switches individually. For more details see the table *inp* in appendix C.

7.1.2 Boundary Conditions and forcing

For the T1 truncation the following lower boundary conditions are specified by external fields: The land sea mask (N002_surf_0172.sra), the surface geopotential (N002_surf_0129.sra) and the surface temperature (N002_surf_0169.sra). The other fields are set by default within the model. Some can be set by namelist parameters (see description of standard model).

The surface fluxes of heat and moisture in the column mode can be influenced by the switch **ZUMIN** in the namelist *fluxpar* which sets the surface wind speed entering the bulk exchange scheme. The default value is set to 1 m/s.

Keeping the standard settings the columns will be forced by the solar forcing corresponding to the grid point where the column is located. For the T1 truncation this mean that the columns are located approximately at gaussian latitudes $\pm 35.26^\circ$. The solar forcing corresponds to the default of the full model. A daily mean insolation is used with an annual cycle. The solar

forcing is also influenced by the climatological ozone distribution which by default also has an annual cycle.

7.2 Graphical User Interface (GUI)

To visualize the time evolution of the column model a vertical Hovmoeller plot has been added to the GUI (picture type: ISOCOL). The sounding device can be also used to visualize the vertical profiles at an arbitrary grid point of the gaussian grid in the full model. By clicking the window the sounding goes from grid point to grid point in meridional direction. The longitude can be selected by the switch **sellon**, which is a parameter of the **inp** namelist in **puma_namelist**. Using the template GUI_sounding.cfg in folder **plasim/dat** the GUI is configured for soundings. In this case **sellon** can be modified in the control window. For more details see chapter 4.

Appendix A

List of Constants and Symbols

Symbol	Definition	Value	Unit
a	earth radius	$6371 \cdot 10^3$	m
A	$= D + \vec{V} \cdot \nabla \ln p_s$		–
\mathcal{A}	absorptivity/emissivity		–
\mathcal{A}_S	surface emissivity		–
$B(T)$	Planck function		Wm^{-2}
cc	cloud cover		–
C_{char}	Charnock constant	0.018	–
C_h	transfer coefficient for heat		–
C_m	drag coefficient for momentum		–
c_p	specific heat of moist air at constant pressure		$\text{J kg}^{-1} \text{K}^{-1}$
c_{pd}	specific heat of dry air at constant pressure	1005.46	$\text{J kg}^{-1} \text{K}^{-1}$
c_{pv}	specific heat of water vapor at constant pressure	1869.46	$\text{J kg}^{-1} \text{K}^{-1}$
c_{pi}	specific heat of sea ice	2070	$\text{W s kg}^{-1} \text{K}^{-1}$
c_{ps}	specific heat of snow	2090	$\text{W s kg}^{-1} \text{K}^{-1}$
c_{pw}	specific heat of sea water	4180	$\text{W s kg}^{-1} \text{K}^{-1}$
c_w	coefficient for the deep ocean heat flux	4	$\text{W m}^{-2} \text{K}^{-1}$
C_w	wetness factor		–
D	scaled divergence		–
E	evaporation		m s^{-1}
E_0	extraterrestrial solar flux density		W m^{-2}
f	Coriolis parameter $=: 2\Omega \sin \varphi$		s^{-1}
F_p	tendency of the first moment $=: \frac{dR_1}{dt}$		$\text{K m}^2 \text{s}^{-1}$
F_q	tendency of the zeroth moment $=: \frac{dR_0}{dt}$		K m s^{-1}
F_q	surface moisture flux		$\text{kg m}^{-2} \text{s}^{-1}$
F_T	surface sensible heat flux		W m^{-2}
F_u	surface zonal wind stress		Pa
F_v	surface meridional wind stress		Pa
F^{LW}	long wave radiation flux density		Wm^{-2}
F^{SW}	short wave radiation flux density		Wm^{-2}
g	gravitational acceleration	9.81	m^{-2}
h_{mix}	mixed layer depth		m
h_{mix_c}	climatological mixed layer depth		m
H_q	effective mixed layer depth $=: \frac{R_0}{T_{mix} - T_{ref}}$		m
H_p	reduced center of gravity $=: \frac{R_1}{R_0}$		m

Symbol	Definition	Value	Unit
J_q	vertical turbulent moisture flux		$\text{kg m}^{-2} \text{s}^{-1}$
J_T	vertical turbulent temperature flux		$\text{K m}^{-2} \text{s}^{-1}$
J_u	vertical turbulent flux of zonal momentum		Pa
J_v	vertical turbulent flux of meridional momentum		Pa
k	von Karman constant	0.4	—
K_h	exchange coefficient for heat		—
K_m	exchange coefficient for momentum		—
L	latent heat		J kg^{-1}
L_f	latent heat of fusion = $L_s - L_v$	$3.28 \cdot 10^5$	J kg^{-1}
l_h	mixing length for heat		m
l_m	mixing length for momentum		m
L_s	latent heat of sublimation	$2.8345 \cdot 10^6$	J kg^{-1}
L_v	latent heat of vapourization	$2.5008 \cdot 10^6$	J kg^{-1}
P_c	convective precipitation		ms^{-1}
P_l	large scale precipitation		ms^{-1}
$P_n^m(\mu)$	associated Legendre function of the first kind		—
p	pressure		Pa
p_S	surface pressure		Pa
p_s	scaled surface pressure		—
q	specific humidity		kg kg^{-1}
Q	total heat flux through sea ice		W m^{-2}
\tilde{Q}	flux correction heat flux through sea ice		W m^{-2}
Q_a	total atmospheric heat flux		W m^{-2}
Q_c	conductive heat flux through sea ice		W m^{-2}
Q_f	heat flux available for freezing sea ice		W m^{-2}
Q_g	heat flux into the soil		W m^{-2}
Q_m	snow melt heat flux		W m^{-2}
Q_o	oceanic heat flux		W m^{-2}
q_S	surface specific humidity		kg kg^{-1}
q_{sat}	saturation specific humidity		kg kg^{-1}
\mathcal{R}	reflexivity/albedo		—
\mathcal{R}_S	surface albedo		—
R_d	gas constant for dry air	287.05	$\text{J kg}^{-1} \text{K}^{-1}$
R_l	surface long wave radiation		W m^{-2}
R_s	surface short wave radiation		W m^{-2}
R_v	gas constant for water vapor	461.51	$\text{J kg}^{-1} \text{K}^{-1}$
R_0	zeroth moment of the temperature distribution		K m
R_1	first moment of the temperature distribution		K m^2
Ri	Richardson number		—
S_w	salinity of sea water	34.7	psu

Symbol	Definition	Value	Unit
t	time		s
\hat{t}	scaled time step		—
\mathcal{T}	transmissivity		—
T	temperature		K
T'	temperature anomaly =: $T - T_0$		—
T_d	deep ocean temperature (at 400m)		K
T_i	sea ice surface temperature		K
T_f	freezing temperature	271.25	K
T_s	surface temperature		K
T_{sea}	sea surface temperature		K
T_{melt}	melting point	273.16	K
T_{mix}	mixed layer temperature		K
T_{mix_c}	climatological mixed layer temperature		K
T_{ref}	asymptotic reference temperature		K
T_w	oceanic temperature profile		K
T_0	reference temperature profile	250.0	K
U	scaled zonal wind =: $u \cdot \cos \varphi$		—
u	zonal wind		m s^{-1}
u_*	friction velocity		m s^{-1}
V	scaled meridional wind =: $v \cdot \cos \varphi$		—
v	meridional wind		m s^{-1}
\vec{v}	horizontal wind vector		m s^{-1}
W_L	cloud liquid water path		gm^2
W_{snow}	mass of snow water		kg
W_{soil}	soil water		m
z	height		m
z_0	roughness length		m
Δt	time increment		s
Δz	height increment		m
α	thermal expansion coefficient $\frac{1}{\rho} \frac{d\rho}{dT}$	$2.41 \cdot 10^{-4}$	K^{-1}
β	back scattering coefficient		—
β_d	diffusivity factor	1.66	—
ζ	scaled vorticity		—
θ	potential temperature		K
κ	R_d/C_{pd}		—
$\bar{\kappa}$	mean heat conductivity in ice and snow		$\text{W m}^{-1} \text{K}^{-1}$
κ_i	heat conductivity in ice	2.03	$\text{W m}^{-1} \text{K}^{-1}$
κ_s	heat conductivity in snow	0.31	$\text{W m}^{-1} \text{K}^{-1}$
λ_h	asymptotic mixing length for heat		m
λ_m	asymptotic mixing length for momentum		m
λ	longitude		—
μ	$\sin \varphi$		—
μ_0	cosine of the solar zenith angle		—
ρ	density of air		kg m^{-3}
ρ_i	density of sea ice	920	kg m^{-3}
ρ_s	density of snow	330	kg m^{-3}
ρ_w	density of sea water	1030	kg m^{-3}
ρ_o	density of fresh water	1000	kg m^{-3}

Symbol	Definition	Value	Unit
σ	normalized pressure coordinate $=: p/p_s$		—
$\dot{\sigma}$	vertical velocity in σ system		—
σ_{SB}	Stefan-Boltzmann constant	$5.67 \cdot 10^{-8}$	$\text{Wm}^{-2}\text{K}^{-4}$
τ_N	cloud optical depth		—
τ_F	time scale for RF		—
τ_R	time scale for NC		—
τ_T	time scale for temperature flux correction		s
τ_h	time scale for depth flux correction		s
ϕ	geopotential height $:= g \cdot z$		$\text{m}^2 \text{s}^{-2}$
$\hat{\phi}$	scaled geopotential height		—
φ	latitude		—
χ	scaled velocity potential		—
ψ	scaled streamfunction		—
Ω	angular velocity of the earth	$7.292 \cdot 10^{-5}$	s^{-1}
$\tilde{\omega}_0$	single scattering albedo		—

Appendix B

Planet Simulator Codes for Variables

Codes available from PUMA-burner (adapted from ECHAM)

Code	Levels	Type	Variable	Unit
110	1	g	mixed layer depth	m
129	1	s	surface geopotential	m ² /s ²
130	NLEV	s	temperature	K
131	NLEV	c	u-velocity	m/s
132	NLEV	c	v-velocity	m/s
133	NLEV	s	specific humidity	kg/kg
135	NLEV	c	vertical velocity	Pa/s
138	NLEV	s	vorticity	1/s
139	1	g	surface temperature	K
140	1	g	soil wetness	m
141	1	g	snow depth (water equi.)	m
142	1	ga	large scale precipitation	m/s
143	1	ga	convective precipitation	m/s
144	1	ga	snow fall	m/s
146	1	ga	surface sensible heat flux	W/m ²
147	1	ga	surface latent heat flux	W/m ²
148	NLEV	c	horizontal streamfunktion	m ² /s
149	NLEV	c	velocity potential	m ² /s
151	1	c	mean sea level pressure	Pa
152	1	s	ln(surface pressure)	
153	NLEV	g	cloud liquid water content	kg/kg
155	NLEV	s	divergence	1/s
156	NLEV	c	geopotential height	gpm
157	NLEV	c	relative humidity	frac.
159	1	g	(u*) ³	(m/s) ³
160	1	ga	surface runoff	m/s

Code	Levels	Type	Variable	Unit
162	NLEV	g	cloud cover	frac.
164	1	ga	total cloud cover	frac.
169	1	ga	surface temperature	K
170	1	g	deep soil temperature	K
172	1	g	land sea mask	[0:sea,1:land]
173	1	g	surface roughness	m
175	1	g	surface albedo	frac.
176	1	ga	surface solar radiation	W/m ²
177	1	ga	surface thermal radiation	W/m ²
178	1	ga	top solar radiation	W/m ²
179	1	ga	top thermal radiation	W/m ²
180	1	ga	u-stress	Pa
181	1	ga	v-stress	Pa
182	1	ga	evaporation	m/s
183	1	g	soil temperature	K
199	1	g	vegetation cover	frac.
203	1	ga	top solar radiation upward	W/m ²
204	1	ga	surface solar radiation upward	W/m ²
205	1	ga	surface thermal radiation upward	W/m ²
207	1	g	soil temperature (level 2)	K
208	1	g	soil temperature (level 3)	K
209	1	g	soil temperature (level 4)	K
210	1	g	sea ice cover	frac.
211	1	g	sea ice thickness	m
212	1	g	forest cover	frac.
218	1	g	snow melt (water equiv.)	m/s
221	1	g	snow depth change (water equiv.)	m/s
230	1	ga	vertical integrated spec. hum.	kg/m ²
232	1	g	glacier cover	frac.

s: PUMA spectral field
g: PUMA grid point field
c: computed by PUMA-burner
a: accumulated

Appendix C

Namelists

C.1 File puma_namelist

C.1.1 Namelist INP

Name	Def.	Type	Description
column	0	int	1: initialize PLASIM for column runs
kick	1	int	0: no noise initialization ($p_s = \text{const.}$) 1: random white noise 2: Equator symmetric random white noise 3: mode (1,2) no random initialization
mars	0	int	1: initialize PLASIM for planet Mars
mpstep	45	int	minutes per step (length of timestep)
nadv	1	int	1: switches horizontal advection on
ncoeff	0	int	spectral coefficients to print in wrspam
ndel(NLEV)	all 2	int	order of hyperdiffusion for each level (2*h)
ndiag	12	int	output interval for diagnostics [timesteps]
ndiagcf	0	int	1: turn on cloud forcing diagnostic
ndiaggp	0	int	1: process franks gp-diagnostic arrays
ndiaggp2d	0	int	number of additional 2-d gp-diagnostic arrays
ndiaggp3d	0	int	number of additional 3-d gp-diagnostic arrays
ndiagsp	0	int	1: process franks sp-diagnostic arrays
ndiagsp2d	0	int	number of additional 2-d sp-diagnostic arrays
ndiagsp3d	0	int	number of additional 3-d sp-diagnostic arrays
ndl(NLEV)	all 0	int	1: activate spectral printouts for this level
neqsig	1	int	1: use equidistant sigma levels
nflux	1	int	1: switches vertical diffusion on
ngui	0	int	1: run with active GUI
nhdiff	15	int	critical wavenumber for horizontal diffusion
nhordif	1	int	1: switches horizontal diffusion on
nkits	3	int	number of short initial timesteps
noutput	1	int	enables (1) or disables (0) output file
npackgp	1	int	1: pack gridpoint fields on output
npacksp	1	int	1: pack spectral fields on output
nperpetual	0	int	radiation day for perpetual integration
nprhor	0	int	1: grid point for print out (only for checks!)
nprint	0	int	1: comprehensive print out (only for checks!)
nrad	1	int	1: switches radiation on
ntime	0	int	1: turn on time use diagnostics
nwpd	1	int	number of writes per day (to puma_output)

Namelist INP continued

Name	Def.	Type	Description
n_days_per_month	30	int	length of month for simple calendar
n_days_per_year	360	int	length of year for simple calendar or 365
n_run_days	-1	int	Simulation time (days to run)
n_run_months	0	int	Simulation time (months to run)
n_run_years	1	int	Simulation time (years to run)
n_start_month	1	int	Starting month
n_start_year	1	int	Starting year
psurf	101100.0	real	global mean surface pressure [Pa]
restim(NLEV)	all 15.0	real	restoration timescale for each level
sigh(NLEV)	all 0.0	real	user definable sigmah array
sellon	0.0	real	longitude of soundings in the GUI
t0(NLEV)	all 250.0	real	reference T_R -temperature profile
tfrc(NLEV)	0,0,0,.. ,1	int	Rayleigh friction timescale τ_F in days
tdissd	0.2	real	diffusion time scale for divergence [days]
tdissq	5.6	real	diffusion time scale for specific humidity [days]
tdisst	5.6	real	diffusion time scale for temperature [days]
tdissz	1.1	real	diffusion time scale for vorticity [days]
time0	0.0	real	start time (for performance estimates)

C.1.2 Namelist PLANET

Name	Def.	Type	Description
akap	0.286	real	kappa
alr	0.0065	real	lapse rate
eccen	0.0	real	eccentricity for fixed orbits
ga	9.81	real	gravity
gascon	287.0	real	gas constant
mvelp	0.0	real	longitude of vernal equinox for fixed orbits (deg)
nfixorb	0	int	1: fix the planetary orbit
obliq	0.0	real	obliquity for fixed orbits (deg)
plarad	6371000.0	real	planetary radius
pnu	0.1	real	time filter
ra1	610.78	real	parameter in Magnus-Teten formula
ra2	17.269	real	parameter in Magnus-Teten formula
ra4	35.86	real	parameter in Magnus-Teten formula
solar_day	86400.0	real	length of solar day
siderial_day	86164.0	real	length of siderial day
ww	7.29e-5	real	$2\pi/siderialday$
yplanet	"Earth"	char	name of planet

C.1.3 Namelist MISCPAR

Name	Def.	Type	Description
nfixer	1	int	1: correct negative moisture
nudge	0	int	1: temperature relaxation in the uppermost level
tnudge	10.0	real	Time scale [d] of the temperature relaxation

C.1.4 Namelist FLUXPAR

Name	Def.	Type	Description
nevap	1	int	1: turn on surface evaporation
nshfl	1	int	1: turn on surface sensible heat flux
nstress	1	int	1: turn on surface wind stress
nvdif	1	int	1: turn on vertical diffusion
vdif_lamm	160.0	real	tuning parameter for vert. diff.
vdif_b	5.0	real	tuning parameter for vert. diff.
vdif_c	5.0	real	tuning parameter for vert. diff.
vdif_d	5.0	real	tuning parameter for vert. diff.
zumin	1.0	real	minimum surface wind speed (m/s)

C.1.5 Namelist RADPAR

Name	Def.	Type	Description
acl2(3)	0.05,0.1,0.2	real	cloud absorptivities spectral range 2
acllwr	0.1	real	mass absorption coefficient for clouds (lwr)
clgray	-1.0	real	cloud grayness
co2	360.0	real	co2 concentration (ppmv)
dawn	0.0	real	zenith angle threshold for night
gsol0	1365.0	real	solar constant (w/m2)
iyrbp	-50	int	Year before present (1950 AD); default = 2000 AD
ndcycle	0	int	switch for daily cycle 1=on/0=off
nlwr	1	int	switch for long wave radiation (dbug) 1=on/0=off
no3	1	int	switch for ozon 1=on/0=off
nrscat	1	int	switch for rayleigh scattering (dbug) 1=on/0=off
nsol	1	int	switch for solar insolation (dbug) 1=on/0=off
nswr	1	int	switch for short wave radiation (dbug) 1=on/0=off
nswrcl	1	int	switch for computed or prescribed cloud props. 1=com/0=pres
rcl1(3)	0.15,0.3,0.6	real	cloud albedos spectral range 1
rcl2(3)	0.15,0.3,0.6	real	cloud albedos spectral range 2
th2oc	0.04	real	absorption coefficient for h2o continuum
tpofmt	1.0	real	tuning of point of mean (lwr) transmissivity in layer
tswr1	0.04	real	tuning of cloud albedo range1
tswr2	0.048	real	tuning of cloud back scattering c. range2
tswr3	0.004	real	tuning of cloud s. scattering alb. range2

C.1.6 Namelist RAINPAR

Name	Def.	Type	Description
clwcrit1	-0.1	real	1st critical vertical velocity for clouds
clwcrit2	0.0	real	2nd critical vertical velocity for clouds
kbetta	1	int	switch for betta in kuo (1/0=yes/no)
ncsurf	1	int	conv. starts from surface (1/0=yes/no)
ndca	1	int	dry convective adjustment (1/0=yes/no)
nmoment	0	int	momentum mixing (1/0=yes/no)
nshallow	0	int	switch for shallow convection (1/0=yes/no)
nprc	1	int	large convective precip (1/0=yes/no)
npri	1	int	switch for large scale precip (1/0=yes/no)
rcrit(NLEV)		real	critical relative hum. for non conv. clouds

C.1.7 Namelist SURFPAR

Name	Def.	Type	Description
noromax	model resolution (NTRU)	int	resolution of orography
nsurf	not active	int	debug switch
oroscale	1.0	real	scaling factor for orography

C.2 File land_namelist

C.2.1 Namelist LANDPAR

Name	Def.	Type	Description
albgmax	0.8	real	max. albedo for glaciers
albgmin	0.6	real	min. albedo for glaciers
albland	0.2	real	albedo for land
albsmax	0.8	real	max. albedo for snow
albsmaxf	0.4	real	max. albedo for snow (with forest)
albsmin	0.4	real	min. albedo for snow
albsminf	0.3	real	min. albedo for snow (with forest)
co2conv	14.0	real	co2 conversion factor
drhsfull	0.4	real	threshold above which drhs=1 [frac. of wsmax]
drhsland	0.25	real	wetness factor land
dsmax	5.00	real	maximum snow depth (m-h20; -1 = no limit)
dsoilz(NLSOIL)		real	soil layer thickness
dwtcini	0.0	real	soil water content (m) for manual initialization (nwtcini=1)
dz0land	2.0	real	roughness length land
dzglac	-1.	real	threshold of orography to be glacier (-1=none)
dztop	0.20	real	thickness of the uppermost soil layer (m)
forgrow	1.0	real	growth factor initialization
gs	1.0	real	stomatal conductance initialization
nbiome	0	int	switch for vegetation model (1/0 : prog./clim)
ncveg	1	int	compute new dcveg (0=keep initial state)
newsurf	0	int	(dtcl,dwcl) 1: update from file, 2:reset
nlandt	1	int	switch for land model (1/0 : prog./clim)
nlandw	1	int	switch for soil model (1/0 : prog./clim)
nwtcini	0	int	switch for manual soil water setting (1/0 : on/off)
rinisoil	0.0	real	soil carbon initialization
riniveg	0.0	real	biomass carbon initialization
rlaigrow	0.5	real	above ground growth factor initialization
rlue	8.0E-10	real	
rnbiocats	0.0	real	
tau_soil	42.0	real	[years] - in landini scaled to seconds
tau_veg	10.0	real	[years] - in landini scaled to seconds
wsmax	WSMAX_EARTH	real	max field capacity of soil water (m)
z0_max	2.0	real	maximum roughness length for vegetation

C.3 File sea_namelist

C.3.1 Namelist SEAPAR

Name	Def.	Type	Description
ncpl_atmos_ice	32	int	atmosphere ice coupling time steps
albsea	0.069	real	albedo for open water
albice	0.7	real	max. albedo for sea ice
dz0sea	$1.5 \cdot 10^{-5}$	real	roughness length sea [m]
dz0ice	$1.0 \cdot 10^{-3}$	real	roughness length ice [m]
drhssea	1.0	real	wetness factor sea
drhsice	1.0	real	wetness factor ice

C.4 File ocean_namelist

C.4.1 Namelist OCEANPAR

Name	Def.	Type	Description
dlayer(NLEV_OCE)	50.0	real	layer depth (m)
ndiag	480	int	diagnostics each ndiag timestep
newsurf	0	int	1: read surface data after restart
nfluko	0	int	switch for flux correction
nocean	1	int	ocean model (1) or climatology (0)
nperpetual_ocean	0	int	perpetual climate conditions (day)
nprhor	0	int	gridpoint for debug printout
nprint	0	int	switch for debug printout
taunc	0.0	real	time scale for newtonian cooling
vdifk	1.0e-4	real	vertikal diffusion coeff. [m**2/s]

C.5 File ice_namelist

C.5.1 Namelist ICEPAR

Name	Def.	Type	Description
newsurf	0	int	1: read surface data after restart
nfluko	0	int	switch for flux correction
nice	1	int	sea ice model (1) or climatology (0)
nout	32	int	model data output every nout time steps
nperpetual_ice	0	int	perpetual climate conditions (day)
nprhor	0	int	gridpoint for debug printout
nprint	0	int	switch for debug printout
nsnow	1	int	allow snow on ice yes/no (1/0)
ntskin	1	int	compute skin temperature (0=clim.
ncpl_ice_ocean	1	int	ice ocean coupling time steps
taunc	0.0	real	time scale for newtonian cooling
xmind	0.1	real	minimal ice thickness (m)