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ICON User's Guide

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Preface

This user guide was assembled and edited based on available documents on the ICON webpage by the persons mentioned at the front page. The content of the user guide follows the requirements of DWD.

Important hints:

In chapter 4 a list of the namelist parameters is given. New and inexperienced users should only modify the namelist parameters that are given in bold letters.

When results produced with ICON are published the following papers have to be cited in the list of references:

?

Information for authors:

Please read the README for further instructions and tamplates. $\,$

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1 Guide for New Users

This tutorial is meant for people with some knowledge and/or experience in modelling and Linux, but which have no experience with the ICON model. In the following we will describe in short how to compile and run ICON on your machine.

1.1 Needed Software

For some components ICON uses external libraries. Therefore you will need some additional software which should be installed on your machine. The following software needed to be installed on your machine:

- NetCDF: NetCDFis a set of software libraries and self-describing, machine-independent data formats that support the creation, access, and sharing of array-oriented scientific data. (Source: http://www.unidata.ucar.edu/software/netcdf/)
- GRIB: GRIB (GRIdded Binary) is a format defined by the WMO (World Meteorological Organization). The use of GRIB in ICON is optional. The ECMWF GRIB API is an application program interface accessible from C, FORTRAN and Python programs developed for encoding and decoding WMO FM-92 GRIB edition 1 and edition 2 messages. A useful set of command line tools is also provided to give quick access to GRIB messages. ICON requires GRIB2 format.

(Source: https://software.ecmwf.int/wiki/display/GRIB/Home)

- MPI: MPI is a library specification for message-passing, proposed as a standard by a broadly based committee of vendors, implementors, and users.
 (Source: http://www.mcs.anl.gov/research/projects/mpi/)
- OpenMP: Jointly defined by a group of major computer hardware and software vendors, the OpenMP API is a portable, scalable model that gives shared-memory parallel programmers a simple and flexible interface for developing parallel applications on platforms ranging from embedded systems and accelerator devices to multicore systems and shared-memory systems. (Source: http://openmp.org/wp/)

1.2 The Source Code

You can obtain the source code on the website of DKRZ:

https://www.dkrz.de/

You can use the following commands to untar the ICON source code:

1.2. THE SOURCE CODE

tar xfvz icon.tar.gz

This will create a folder icon-1.0 inside your current directory. Within the ICON User Guide, this folder will further on be called \$ICONDIR.

1.2.1 Directory structure

Within \$ICONDIR, you will find a set of subdirectories. The important subdirectories are described in the following.

build

Within the \$ICONDIR/build directory, a subdirectory with the name of your computer architecture is created at compilation. Within this subdirectory, a bin subdirectory containing the binary control_model and several further subdirectories containing the compiled module files are created at compilation.

config

Inside the \$ICONDIR/config directory, different machine dependent configuration are stored within the configuration files. You can find a description of how to use and set up such configuration files in chapter 1.3.

data

Within the \$ICONDIR/data directory, you will find divers input datasets. For example, there are the datasets "rrtmg_lw.nc" and "ECHAM6_CldOptProps.nc", which are necessary for the radiation scheme (see sec. 1.6.1).

doc

Within the \$ICONDIR/doc directory, several documentations for ICON are stored. There are according subdirectories for scientific (\$ICONDIR/doc/science), technical (\$ICONDIR/doc/technical) and programming style guides (\$ICONDIR/doc/style).

externals

Within the \$ICONDIR/externals directory, external libraries for ICON are stored. Currently, it is the mtime library which is used to convert different date time formats.

include

Within the \$ICONDIR/include directory, interfaces to libraries needed by ICON are stored. Currently, the interface to the CDI library is stored inside this directory.

run

Within the \$ICONDIR/run directory, namelist descriptor files as well as the full namelist documentation are stored. The namelist descriptor files can be used to generate runscripts. Further information can be found in 1.6.

src

Within the \$ICONDIR/src directory, the source code of ICON including the main program and ICON modules can be found. The modules are ordered in several subdirectories which are described in the following.

The main program control_model.f90 can be found inside the subdirectory \$ICONDIR/src/drivers. Additionally, this directory contains the modules for a hydrostatic and a nonhydrostatic setup.

The configuration of an ICON run is performed within the modules inside \$ICONDIR/src/configure_model and \$ICONDIR/src/namelists. Modules regarding the configuration of idealized test cases can be found inside \$ICONDIR/src/testcases.

The dynamics of ICON are inside \$ICONDIR/src/atm_dyn_iconam and the physical parameterizations inside \$ICONDIR/src/atm_phy_nwp. Parameterizations for the interactions with the surface can be found inside \$ICONDIR/src/lnd_phy_nwp.

Shared infrastructure modules for 3-D and 4-D variables can be found within \$ICONDIR/src/shared. The according routines for 2-D fields (e.g. external parameters) are stored within \$ICONDIR/src/shr_horizontal.

Modules handling the parallelization can be found in \$ICONDIR/src/parallel_infrastructure.

Input and output modules are stored in \$ICONDIR/src/io.

The modules for the grid generator, as described in chapter 1.8 can be found inside \$ICONDIR/src/grid_generator.

support

Within the \$ICONDIR/support directory, the CDI library is stored.

vertical_coord_table

Inside the \$ICONDIR/vertical_coord_tables directory, information files describing the relation between model layer and height are stored.

1.3 Configuration and Compilation

To ease up the compilation a configure-file is provided which should take over the main work. This Autoconf configuration is used to analyze the computer architecture (hardware and software) and set user specified preferences, e.g. the compiler. This preferences are read from config/mh-<OS>, where <OS> is the identified operating system. Operating systems are listed in the configure-files in \$ICONDIR/config/ with the according files mh-<OS>. If your machine is not listed you can add a config-file with your own <OS> based on the given mh-<OS> files. If different compilers are available, the mh-<OS> file may contain a case construct to distinguish them. If your <OS> is not recognized but is one of the listed <OS> you can invoke the configure file with the according option --host=\$HOST. Examples for the DWD CRAY system are given in the boxes.

1.3.1 Description of the Configuration Files

To add a specific compiler or change your compiler flags, you have to enter the \$ICONDIR/config/mh-<OS> according to your operating system <OS>. For the DWD CRAY, the compiler flags in mh-linux look like the following:

```
CRAY EXAMPLE: Compiler Flags inside mh-linux
    config_compiler=cray
        CC
                    = cc
    FC
                = ftn
                = "$FC"
   F77
   FFLAGS
                = -v -D__LOOP_EXCHANGE -D__MIXED_PRECISION -Df2cFortran
-e Z -em -hflex_mp=conservative -hfp1 -hadd_paren -r am -Ktrap=divz,ovf
    CFLAGS
                = -I${GRIB_API}/include -v -Df2cFortran
-DHAVE_CF_INTERFACE -DHAVE_LIBNETCDF -DHAVE_LIBGRIB
-DHAVE_LIBGRIB_API -O3 -D__SVN_VERSION="${SVNVERSION}"
                = "$FFLAGS"
    F77FLAGS
                = "-v"
    FCLIBS
    GEN_FLAGS
    FDEBUG
                = -g -R abc
    OMPFLAG
                = -mp
    DEFOPT
                = -D
                = -D
   DEFCOPT
   MODOPT
                = -I
   MODDIR
    ;;
```

The cray) in this example gives the name of this specific configuration. It can be addressed by a flag at configuration. For this example, the according command to choose this setting would

be ./configure --with-fortran=cray (see section 1.3.2). Like this, you can create your own configuration by adding a new compiler.

CC, FC and F77 are the compiler directives for C-Compiler, FORTRAN2003-Compiler and FORTRAN77-Compiler. The according compiler flags are set via CFLAGS, FFLAGS and F77FLAGS. The variable to set an OpenMP flag is called OMPFLAG. Libraries are set via FCLIBS.

1.3.2 Configuring and Compiling the Code

To configure the source code go to \$ICONDIR and give:

```
./configure
./build_command
```

If you want to use another compiler than the default compiler you give:

```
./configure --with-fortran=<compiler>
./build_command
```

where <compiler> is {gcc,nag,intel,pgi,cray}.

```
CRAY EXAMPLE: Configure + Make
./configure --with-fortran=cray}
./build_command
```

Note, that CRAY compiler environment (cce) versions 8.2.x do not work with ICON. The CRAY configuration is expanded to the following:

```
CRAY EXAMPLE: Configuration

ftn -I../module -v -D__LOOP_EXCHANGE -D__MIXED_PRECISION -Df2cFortran -e

Z -em -hflex_mp=conservative -hfp1 -hadd_paren -r am -Ktrap=divz,ovf

-D__ICON__ <object files> -L/usr/local/pkg/grib_api/1.11.0/CRAY/lib

-L../lib -lsupport -lgrib_api_f90 -lgrib_api -lmtime $(LAPACK_LIB)

$(NETCDF_LIB) $(HDF5_LIB) $(SZIP_LIB) $(ZLIB_LIB) $(MPI_LIB)

$(PROFILE_LIB) $(SCT_LIB)
```

ICON is parallelized using MPI and OpenMP. You can control the parallelization to be used by giving:

```
./configure --with-mpi/--without-mpi --with-openmp/--without-openmp ./build_command
```

By default the options are set to --with-mpi --without-openmp. After a successful build, you will find the ICON executable named control_model inside \$ICONDIR/build/<0S>/bin/. The CRAY

Fortran compiler is an exception, as the command includes automatically OpenMP. Therefore, although selecting –without-openmp, OpenMP is used.

If you wish to re-configure ICON it is advisable first to clean the old setup by giving:

make distclean

Some more details on configure options can be found in the help of the configure command:

./configure --help

1.4 Running the model (Test scripts for ICON)

1.4.1 Principles of testing ICON

The ICON developers use the buildbot tool in order to perform automated tests on selected ICON experiments at regular time intervals. The buildbot tool launches the respective test scripts on various computer platforms and documents success or failure on a special web site (https://buildbot.zmaw.de/icon The automated tests are performed on the newest model revision as available in the ICON repository. Furthermore, tests can be "forced", i.e. started by hand, at any time specifying a certain experiment and model revision of any branch of the repository. However, it is impossible to test two revisions against each other or to test a local revision. Here, we present new tests for ICON into which various experiments are integrated and which are designed such that they can either be used in the framework of the buildbot tool or be started manually without reference to build-bot (e.g. on a PC at MPI Hamburg). The tests are designed to trap certain technical errors and comprise the following experiments:

base test: Just a simple base run over a short time period for a specific experiment. This run will be called simulation A in the following.

update test: In addition to the model revision to test, the so-called "test revision", a "reference revision" can be specified. A short simulation A of the test revision over one hour is performed during which restart files are written. The same simulation is performed for the reference revision (simulation A'). The "update test" is said to be passed if there are no differences in the output of simulation A and A' using the "cdo diff" command on the time steps in the output specified by the user.

restart test: In addition to a base simulation A, a second simulation B restarts ICON at some time after the initial date. The "restart test" is said to be passed if there are no differences in the output for the time steps after the restart between the original and the restarted simulation using the "cdo diff" command.

nproma test: The nproma test performs a base simulation A and a simulation C with a different value of nproma. Instead of nproma of simulation A, a value of 17 is used in simulation C or, if nproma = 17 for simulation A, nproma = 19 is used for simulation C. The nproma test is said to be passed if the "cdo diff" command does not find differences in the output.

mpi parallel test: The mpi parallel test performs a base simulation A and a simulation D with a reduced number of MPI threads compared to the base simulation A. If more than one threads are used on each node, the number of threads on each node is reduced by one. If only one thread is used on each node, the number of nodes is reduced by one. If only one process is used, no mpi parallel test is performed. The parallel test is said to be passed if the "cdo diff" command does not find differences between the output files.

openmp parallel test: The openmp parallel test performs a base simulation A and a simulation E with a reduced number of openmp threads compared to the base simulation A. If only one openmp thread was used, no openmp test is performed. The openmp parallel test is said to be passed if the "cdo diff" command does not find differences between the output files.

The testing procedure is such that tests can be combined. Furthermore, the test script can be asked to re—use existing runs without repeating these runs.

Only the following experiments are included into the test script:

atm_amip_test: Non–hydrostatic AMIP–like simulation but with transient solar irradiance using ECHAM physics.

atm_icoles_nested: Nonhydrostatic atmosphere only simulation with a regional grid refinement.

atm_jww_hs_test: Jablonowski Williamson baroclinic wave test for a hydrostatic atmosphere.

oce_omip_0160km: Ocean only experiment with a 160km resolution.

1.4.2 Description of test script

The test script icon-dev.checksuite is located in the run/checksuite.icon-dev directory of ICON and uses the following run script of the run directory for the experiments: (i) exp.atm_amip_test for the AMIP-type experiment atm_amip_test, (ii) exp.atm_icoles_nested for the atmosphere experiment with a grid refinement, (iii) exp.atm_jww_hs_test for the Jablonowski Williamson baroclinic wave test, and (iv) exp.oce_omip_0160km for the ocean only experiment oce_omip_0160km.

These run scripts contain all necessary namelist groups and links to files that contain the initial and boundary conditions. By the standard make_runscripts command invoked inside icon-dev.checksuite, this script is transformed into the actually used form that contains an additional suffix .run at the end of its name. Attention: icon-dev.checksuite generates the specific run script by default and overwrites those that are present. The script can be forced to use present runscripts. For the various test runs for each experiment, these .run files are copied and edited by sed.

Here follows a more detailed description of the script:

icon-dev.checksuite: This script uses the make_runscripts command to produce exp.<exp_name>.run from the basic run script exp.<exp_name>. The default is that any existing run script is overwritten but there is an option to keep existing runscripts. These run scripts are then modifed by sed commands in order to perform the various test runs. Once the test runs are finished, the function diff_results of icon-dev.checksuite is called to determine the differences between those runs.

exp.<experiment>: These scripts contain all settings for the base simulation in one experiment. To date, the experiments <experiment> = atm_amip_test, atm_icoles_nested,

atm_jww_hs_test, and oce_omip_160km can be used in the tests. The base script exp.<experiment> will be transformed by make_runscripts into a script that can actually run the ICON model. The resulting exp.<experiment>.run scripts will then be copied to exp.<experiment>_base.run, exp.<experiment>_restart.run, exp.<experiment>_nproma.run, exp.<experiment>_mpi.run, and exp.<experiment>_omp.run to perform simulations A, B, C, D, and E, described in section 1.4.1, respectively. The latter scripts are then modified by icon-dev.checksuite using sed according to the needs of the respective runs.

diff_results. This function compares two simulations. The five arguments contain the base path of the model (.../icon-dev/ for example), and the name of the experiment to be compared (e.g. <experiment>_base) for the two experiments, respectively. The path of the models can be identical (e.g. for the restart or nproma tests that are performed on the same model revision). The fifth argument is the name of the test (update, restart, nproma, mpi, omp) and is only used to produce more legible output. However, the diff_results function needs further information that is provided by variables that are set in the main script: (i) the respective infix of the output files in variable TYPES (e.g. atm_phy), the output dates and time in variable DATES (e.g. 19780101T004000Z) as they figure on the output filenames, and the restart date in RESTART_DATE. These three variables can be set as arguments to the options -t, -d, and -s in a call to icon-dev.checksuite, respectively. The diff_results function checks for differences between two experiments by the cdo diff command. If the variable SUB_FILES is set to 'yes', e.g. by the use of the -u option in the call of icon-dev.checksuite, the variables of the respective outputfiles are subtracted from each other resulting in difference files

```
diff_<EXP2>_<TYPE>_<DATE>-<EXP1>_<TYPE>_<DATE>.nc
for <TYPE> in TYPES and <EXP[12]> one of <experiment>_base,
<experiment>_restart, <experiment>_nproma, <experiment>_mpi,
<experiment>_openmp, or <experiment>_update. The difference files are written to
the path of experiment EXP2.
```

1.4.3 Usage

There are three different ways to use the "check suite":

- (i) Start on the command line: The test script icon-dev.checksuite can be called on the command line from the run/checksuite.icon-dev directory. All the below described options are available on the command line and the full functionality can be used via the command line options easily.
- (ii) Submit to queue: Like buildbot does, it is possible to run make_runscripts on a respective test experiment script located in icon_dev/run and to submit the resulting run script to the respective queuing system. E.g. from exp.test_atm_amip, the runscript exp.test_atm_amip.run is generated and can be submitted. exp.test_atm_amip is just a link to checksuite.icon-dev/check.atm_amip. In order to use the full functionality of icon-dev.checksuite, various environment variables have to be set in exp.test_atm_amip. This way of calling run/checksuite.icon-dev is good for testing on computers with a queuing system. To date, the exp.test_atm_amip and exp.check_oce_omip_160km are the only test scripts that are available.
- (iii) Buildbot: The script calling icon-dev.checksuit can be used by buildbot. In this case, it is

important to check that the correct values of all the environment variables are set in the run scripts mentioned in paragraph (ii).

The calling syntax of icon-dev.checksuite is:

Description of options:

- -c: colour line output. Colour output should not be used when the script is called by buildbot.
- -d: dates for which outputfiles exist. The default depends on the experiment.
- -e: experiment on which tests have to be performed. Currently, the non-hydrostatic amip-like experiment atm_amip_test, the atmospheric experiment including a grid refinement atm_icoles_0160km, the Jablonowski Williamson baroclinic wave test on a hydrostatic atmosphere atm_jww_hs_test, and the ocean only experiment oce_omip_0160km are supported.
- -f: The argument yes forces to create run scripts even if they already exist (default), no creates run scripts only if they are not yet present.
- -h: display help
- -m: describes the test mode by its arguments that are one of b(ase), u(pdate), r(estart), n(proma), m(pi), o(mp), ur, un, um, uo, rn, rm, ro, nm, no, mo, urn, urm, uro, unm, uno, umo, rnm, rno, rmo, nmo, urnm, urno, urmo, urmo, urnmo, urnmo. The first five tests modes describe the sole base run, or the update—, restart—, nproma—, mpi—, and omp—tests, respectively. The last 26 acronyms describe combined tests where each single test is represented by its initial letter. The default test mode is rnmo.
- -o: The argument of this option can be either yes or no depending on whether existing test simulations shall be overwritten (-o yes) or will be re—used for the current tests (-o no). The default is -o yes, so all existing experiments are automatically overwritten if not specified otherwise.
- -r: The argument of this option gives the absolute path to the reference model. If the test mode includes an update test, it is mandatory. No default.
- -s: Restart date for the restart test as given by the time settings in the respective experiment.

 Default depends on the experiment.
- -t: "Types" (infixes) of output files that have to be compared. The infixes depend on the experiment and are set by default accordingly.
- -u: If files in the various test runs differ, calculate the difference by cdo sub.

Corresponding to the options on the command line, the following environment variables can be set in the exp.test_<experiment>:

```
-c: COLOUR='yes'|'no'. Not recommended in use with buildbot.
-d: DATES=<date_string>.
-e: EXPERIMENT=<name>.
-f: FORCE_MRS='yes'|'no'
-m: MD=<test_mode>
-o: OVERWRITE='yes'|'no'
-r: REFERENCE=<reference_model_path>
-s: RESTART_DATE=<restart_date>
-t: TYPES=<file_type_infixes>
-u: SUB_FILES='yes'|'no'
```

1.4.4 Examples

```
icon-dev.checksuite -o no -c -u -e oce_omip_160km
```

This command runs the rnmo, i.e. the restart, nproma, mpi and openmp test on the experiment oce_omip_160km. Existing runs are not overwritten (-o no), there is colour output (-c), and the difference files are calculated between the various test experiments and the base run (-u).

```
icon-dev.checksuite -c -f no -m ur -r <path>
```

This command performs the update and restart test (-m ur) on the atm_amip_test experiment, colour output ist switched on (-c), the reference model is given in <path>, the runscripts are not newly generated if they are already there (-f no).

1.5 Running the Model (Idealized Cases)

To shed light on the functionality and the quality of the dynamical core, setups for two test cases are presented in the following. Additionally, results of these test cases are shown. These tests are classified in short deterministic test cases (typically a simulation period of about 10-30 days) and tests in a climate mode (typically a multi-year period). This section concentrates on the first class, which starts from prescribed initial conditions (ideally provided in analytic form). The simulation results are either compared to analytic solutions (if available) or high-resolution reference solutions. For a list of available testcases, the reader is referred to the namelist section (4).

1.5.1 Jablonowski-Williamson test

The Jablonowski-Williamson Test (?) is a standard test for dynamical cores in global models and can be run for dry dynamics only - as it is intended for- but full physics can be also tested.

Input von Daniel Reinert is expected here.

Setup

For full physics, two additional namelist parameters are introduced in the testcase_nml to control the initial moisture in the atmosphere:

- Here rh_at_1000hpa to be set between 0 and 1. The default is set to 0.7 which gives a quite smooth start. If you really want to see early onsets of convection and microphysics you have to tune this parameter.
- qv_max is usually set to 20.e 3kg/kg and refers to the maximum value in the tropics.

Input Data

GRID

Results

The Jablonowski-Williamson steady-state test is based on a zonally symmetric, strongly baroclinic atmosphere. Initially, it is in a hydrostatic and geostrophic balance and therefore should remain stationary if no perturbation is imposed. Grid irregularities can disturb this stationary conditions and hence the test identifies the presence and magnitude of grid imprinting of a numerical model. For the Jablonowski-Williamson baroclinic wave test, a weak (and unbalanced) perturbation disturbs the initial wind. This test highlights the diffusivity (or effective resolution) of a dynamical core and the presence of phase speed errors in the advection of poorly resolved structures.

1.5.2 Mountain induced Rossby waves

In order to test the model dynamics in dry stage but with real or any complex topography one can choose the mountain induced Rossby wave test case and select different types of topography. The following namelist parameters give an example how to perform such an idealized simulation.

```
NAMELIST EXAMPLE for moutain induced Rossby waves
! nh_testcase_nml: idealized testcase specification
&nh_testcase_nml
nh_test_name
                        = 'mrw_nh' ! testcase selection
                        = 20.0
                                   ! initial u-component
u0_mrw
mount_height_mrw
                        = 2000.0
                                   ! maximum mountain height
mount_half_width
                        = 1.5e06
                                   ! half width of mountain
mount_longctr_mrw_deg = 90.0
                                   ! longitude: center of the mountain
mount_latctr_mrw_deg
                        = 30.0
                                   ! latitude : center of the mountain
! run_nml: general switches
&run nml
ltestcase
          = .TRUE.
                      ! idealized testcase runs
```

```
num_lev
           = 90
                     ! number of full levels (atm.) for each domain
lvert_nest = .TRUE.
                     ! vertical nesting
nsteps
           = 1000
                     ! number of time steps of this run
dtime
           = 288
                     ! timestep in seconds
                     ! compute adiabatic dynamic tendencies
ldynamics = .TRUE.
ltransport = .FALSE. ! compute large-scale tracer transport
ntracer
                     ! number of advected tracers
           = 0
iforcing
           = 0
                     ! forcing by parameterized processes
msg_level = 7
                     ! controls printout during runtime
ltimer
           = .FALSE. !monitoring the runtime of specific routines
output
           = "nml"
                     ! main switch for components of the model output
```

Initial conditions

Applying this namelist parameters the topography shown in Fig. 1.1 is used.

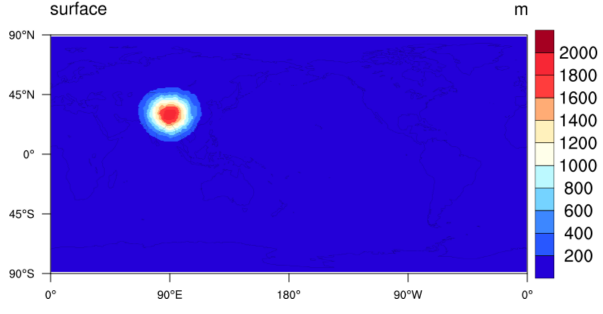


Figure 1.1: Topography of the test case

The v-component of the wind speed is is initialized with zero at all grid points, the initial conditions for the u-component are shown in Fig. 1.2.

Results after 16 days

The u-component after sixteen days of simulation at 700 hPa is shown in Fig. 1.3, the corresponding v-component is shown in Fig. 1.4, and Fig. 1.5 shows the vorticity.

Input Data

With the exception of the grid file no further input files are necessary.

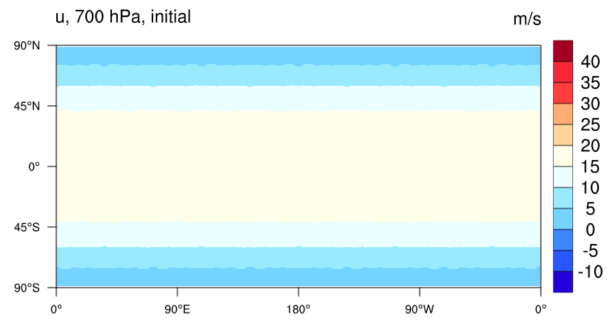


Figure 1.2: Spatial distribution of the initialized u-component at 700 hPa

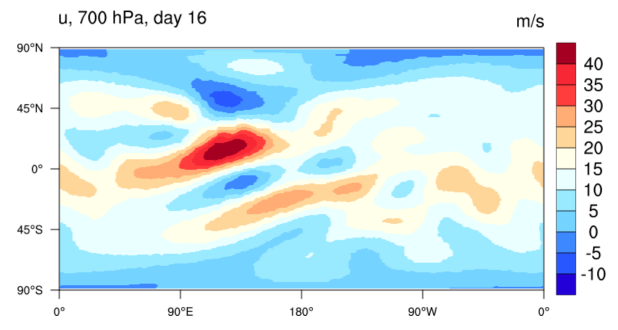


Figure 1.3: Spatial distribution of u-component after 16 days of simulation at 700 hPa

1.5.3 cr2016_02_16_rjs: Aqua planet in ICON

The Aqua Planet Experiment (APE) is incorporated in ICON as a set of testcases, all of them having in common that the bottom boundary consists of water only, with all land surface and orography being removed. Originally APEs where constructed as a test-bed for model intercomparison regarding the interaction of dynamics and physical parameterizations in atmospheric General Circulation Models (GCMs) ??. However, this idealized setup is also well suited for studies focusing on the interaction of basic atmospheric phenomena and the ocean surface, e.g. tropical convection, when a realistic representation of the atmosphere is secondary.

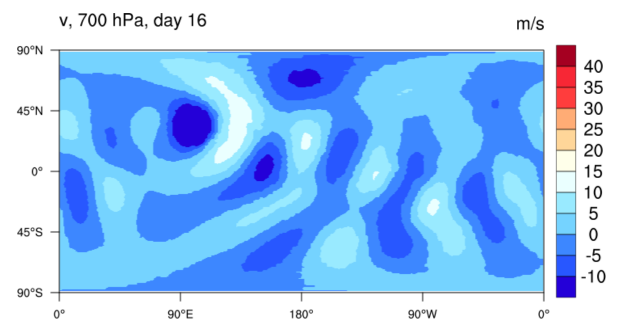


Figure 1.4: Spatial distribution of v-component after 16 days of simulation at 700 hPa

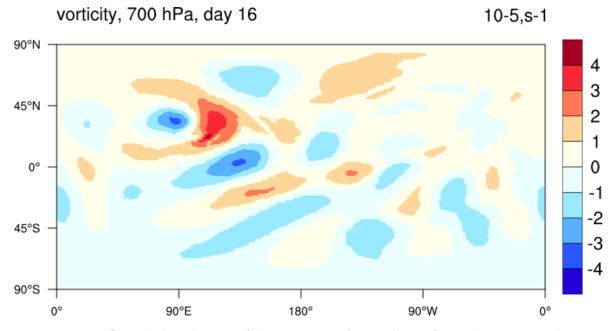


Figure 1.5: Spatial distribution of the vorticity after 16 days of simulation at 700 hPa

The APE is one of the test cases of ICON and can be switched on by setting the respective nh_test_name variable in the nh_testcase_nml namelist, see Listing 1.1.

Sea Surface Temperatures (SSTs) can either be prescribed by one of several analytical functions and kept constant in time, or alternatively SSTs may interact with the atmosphere by the use of a slab ocean layer. The slab ocean is not tested yet. The following SSTs $T_{\rm surf}$ are available from mo_ape_params.f90 under the respective keywords ($T_{\rm m}=273.15\,{\rm K}$):

keyword sst1

$$T_{\text{surf}} = \begin{cases} T_{\text{m}} + 27 \,\text{K} \left(1 - \left(\sin(3\phi/2) \right)^2 \right) & \text{for all } -\pi/3 < \phi < \pi/3 \\ T_{\text{m}} & \text{otherwise} \end{cases}$$
(1.1)

keyword sst2

$$T_{\text{surf}} = \begin{cases} T_{\text{m}} + 27 \,\text{K} \left(1 - \frac{3}{\pi} |\phi|\right) & \text{for all } -\pi/3 < \phi < \pi/3 \\ T_{\text{m}} & \text{otherwise} \end{cases}$$
(1.2)

keyword sst3

$$T_{\text{surf}} = \begin{cases} T_{\text{m}} + 27 \,\text{K} \left(1 - \left(\sin(3\phi/2) \right)^4 \right) & \text{for all } -\pi/3 < \phi < \pi/3 \\ T_{\text{m}} & \text{otherwise} \end{cases}$$
(1.3)

keyword sst4: This distribution has its temperature peak at 5°N:

$$T_{\text{surf}} = \begin{cases} T_{\text{m}} + 27 \,\text{K} \left(1 - \left(\sin(1.6363(\phi - \frac{\pi}{36})) \right)^2 \right) & \text{for all} \quad \pi/36 < \phi < \pi/3 \\ T_{\text{m}} + 27 \,\text{K} \left(1 - \left(\sin(1.3846(\phi - \frac{\pi}{36})) \right)^2 \right) & \text{for all} \quad -\pi/3 < \phi \le \pi/36 \end{cases}$$
 (1.4)
$$T_{\text{m}}$$
 otherwise

keyword sst_qobs

$$T_{\text{surf}} = \begin{cases} T_{\text{m}} + \frac{27}{2} \,\text{K} \left(2 - (\sin(3\phi/2))^2 \left(1 + (\sin(3\phi/2))^2 \right) \right) & \text{for all } -\pi/3 < \phi < \pi/3 \\ T_{\text{m}} & \text{otherwise} \end{cases}$$
(1.5)

keyword sst_ice

$$T_{\text{surf}} = \begin{cases} T_{\text{m}} + 1.9 \,\text{K} + 27 \,\text{K} \left(1 - \left(\sin(3\phi/2) \right)^2 \right) & \text{for all } -\pi/3 < \phi < \pi/3 \\ T_{\text{m}} - 1.9 \,\text{K} & \text{otherwise} \end{cases}$$
(1.6)

keyword sst_const

$$T_{\text{surf}} = T_{\text{m}} + 29 \,\text{K} \tag{1.7}$$

The various SSTs can be set in the namelist nh_testcase_nml by attributing the corresponding keyword to the variable ape_sst_case. For an example see Listing 1.1.

Listing 1.1: Testcase namelist for APE (exp.atm_ape_test)

The APE can be conducted with the MPI/ECHAM physics only since the use of the PSRAD radiation is essential for the APE. A standard example script exp.atm_ape_test is provided. In this standard version, a Kepler orbit with no eccentricity and zero obliquity is used. The orbit parameters are set in the psrad_orbit_nml namelist (Listing 1.2).

Listing 1.2: Orbit namelist for APE (exp.atm_ape_test)

```
&psrad_orbit_nml
cecc = 0.0     ! zero excentricity
cobld = 0.0     ! zero obliquity
l_orbvsop87 = .FALSE.     ! use Kepler orbit instead of historic one
/
```

If a circular orbit with no obliquity is used, a year with months of a different number of days does not make sense. Instead, a 360 day year should be used. The calendar must be set by two variables at the beginning of the exp.atm_ape_test script (Listing 1.9).

Listing 1.3: Calendar for APE (exp.atm_ape_test)

```
calendar="'360udayuyear'"
calendar_type=2
```

The solar irradiation is set to the preindustrial value, i.e. to a value of $1360.875 \,\mathrm{W/m^2}$. The APE uses a composition of the atmosphere where the O_3 concentration is read from a special file. Prognostic water vapour and prognostic liquid and ice cloud water is used together with constant CO_2 in space and time, and no aerosols. All other greenhouse gases are set to zero (Listing 1.4).

Listing 1.4: Solar irradiation and composition of atmosphere (exp.atm_ape_test)

```
&radiation_nml
            = 2 ! preindustrial solar constant of 1360.875 W/m^ at
isolrad
    1 AE
irad_h2o = 1 ! prognostic vapor, liquid and ice
irad_co2
            = 2 ! constant co2
irad_o3
            = 4 ! perpetual january of ozone file linked
irad_ch4
            = 0 ! no ch4
irad_n2o
            = 0 ! no n2o
irad_o2
           = 0 ! no o2
irad_cfc11 = 0 ! no cfc11
irad_cfc12 = 0 ! no cfc12
irad_aero = 0 ! no aerosol
```

The O₃ concentrations are read from the file ape_o3_iconR2B04-ocean_aqua_planet.nc that is provided in the directory data/external/ape_ozone of the ICON main directory. Table 1.1 lists possible ozone files.

1.5.4 cr2016_02_16_rjs: Radiative—Convective equilibrium in ICON using ECHAM physics

The radiative–convective equilibrium (RCE) offers a possibility to improve our fundamental understanding of processes in the atmosphere and their impact on climate change (e.g. ?). The idea behind this simplified modeling of the atmosphere is that the basic atmospheric structure, especially in the tropics, is determined by the balance between cooling of the atmosphere through radiative processes and a commensurate heating through convection, mainly by the net release of latent heat through precipitation.

file name description ape_o3_iconR2B03-ocean_aqua_planet.nc contains one time step of ozone values, interpolated from ape_o3_T42_1Pa.nc to r2b3 resolution ape_o3_iconR2B04-ocean_aqua_planet.nc contains one time step of ozone values, interpolated from ape_o3_T42_1Pa.nc to r2b4 resolution contains the same ozone values as the ape_o3_R2B04_1Pa_spr0.90-cell.nc file above ape_o3_R2B04_hex_1Pa_c.nc unclear, contains 10242 grid points only instead of 20480 as the files for r2b4 resolution (hexagons instead of triangles?) tor120000s64_ozone_CMIP5_aqua_1870_march.nc one time step on 8192 cells, 36 levels, seems to be on a torus grid.

Table 1.1: Possible ozone file

The RCE has been investigated in models of different complexity, ranging from simple energy balance, 1-dimensional column models to high resolution LES simulations. The RCE is also implemented into the general circulation model ICON by creating a model configuration, where the resulting climate is given merely through the balance of radiative processes and convection. Columns can interact with each other and thus create a mean three-dimensional circulation which develops interactively, although it is very different from the general circulation we know from the real Earth. E.g., the RCE results in slowly moving convective clusters of sometimes continental extension (?).

To inhibit net energy transport from the tropics to the poles, homogeneous boundary conditions are specified, where every gridpoint of the sphere receives the same incoming solar radiation (e.g. about $340\,\mathrm{W/m^2}$). A diurnal cycle may be switched on, but is kept exactly the same for each column representing a pulsating light source shining from all directions equally. The Earth's rotation velocity is set to zero. In the standard RCE configuration, land—sea contrasts are removed by specifying an underlying mixed—layer ocean with a constant ocean albedo, but can easily be included in idealized form for land—sea contrast studies (?). This model version has not been tested for possible equilibria dependence on the initial boundary conditions yet, nor for complete isotropy of variables expected from the homogeneous boundary conditions.

Setting initial and boundary conditions and parameters for the RCE

We describe the settings in the standard example script exp.atm_rce_test here. In that case, the mixed layer ocean is switched on by

Listing 1.5: Mixed layer ocean switch in exp.atm_rce_test

```
&echam_phy_nml
...
lmlo = .TRUE.
...
/
```

For the initialization of the RCE configuration, we use the nh_testcase_nml namelist.

Listing 1.6: Testcase namelist for RCE in exp.atm_ape_test

```
&nh_testcase_nml
nh_test_name = 'RCE_glb'
ape_sst_case = 'sst_const'
ape_sst_val = 25.
tpe_temp = 298.15
tpe_psfc = 1013.25e2
//
```

In namelist nh_testcase_nml (Listing 1.6), the variable nh_test_name set to 'RCE_glb' has the effect to initialize a global model with all values not depending on the geographical position. The primary initialization is done in init_nh_state_rce_glb of module mo_nh_rce_exp.f90. After this primary inizialization, a random noise is added to the variables. The variable ape_sst_case is used to initialize the ocean surface temperature by a constant throughout the globe whereas ape_sst_val gives the value of the sea surface temperature in degrees centigrade. The variable tpe_temp determines the temperature of the atmosphere in Kelvin that is independent of altitude at the beginning. The variable tpe_psfc gives the surface pressure in Pascal.

The RCE model can be run in a configuration with the MPI/ECHAM physics only, since the use of the PSRAD radiation is essential for the special radiation settings of the RCE. A uniform irradiation of the globe that may undergo a daily cycle is switched on by setting <code>l_sph_symm_irr</code> to .TRUE. in namelist <code>psrad_orbit_nml</code> (see Listing 1.7). However, a Kepler orbit with no eccentricity has to be used to warrant a sun—earth distance that is constant in time. This is assured by the three first variables in the namelist <code>psrad_orbit_nml</code> as shown in Listing 1.7. The obliquity is set to zero for convenience.

Listing 1.7: Orbit namelist psrad_orbit_nml for RCE in exp.atm_rce_test

```
&psrad_orbit_nml
cecc = 0.0
cobld = 0.0
l_orbvsop87 = .FALSE.
l_sph_symm_irr = .TRUE.
//
```

The angular velocity of the earth is set to zero in order to exclude any quantity that depends on latitude as the Coriolis force. To this end, grid_angular_velocity is set to zero in namelist grid_nml (see Listing 1.8).

Listing 1.8: Grid namelist grid_nml for RCE in exp.atm_rce_test

```
&grid_nml
  grid_angular_velocity = 0.
/
```

The diurnal cycle can be switched on and off with the variable ldiurn of namelist radiation_nml (see Listing 1.10).

If a circular orbit with no obliquity is used, a year with months of a different number of days does not make sense. Instead, a 360 day year should be used. The calendar must be set by two variables at the beginning of the exp.atm_ape_test script (Listing 1.9).

Listing 1.9: Calendar for APE (exp.atm_ape_test)

```
calendar="'360_day_year'"
calendar_type=2
```

The solar irradiation has to be set to an average value that corresponds to the actual energy flux into the atmosphere of the real earth. If spherically symmetric irradiation without diurnal cycle is chosen, isolrad has to be set to 5, if the diurnal cycle is switched on, isolrad has to be set to 4, respectively (see Listing 1.10). These settings result in a global mean insolation of $340.3 \,\mathrm{W/m^2}$. However, the sum of the 14 solar wavelength bands is higher $(433.3371 \,\mathrm{W/m^2}$ and $1069.315 \,\mathrm{W/m^2}$ for isolrad = 5, 4, respectively), due to the applied solar zenith angles and the eventual diurnal cycle.

The composition of the atmosphere used in the RCE model is described by the corresponding switches in namelist radiation_nml (see Listing 1.10). The O₃ concentration is read from a special file r2b4_ozone_rce.nc that is distributed with the program code in the directory data/external/ape_ozone of the ICON main directory. Prognostic water vapour and prognostic liquid and ice cloud water is used together with constant CO₂ in space and time, and no aerosols. All other greenhouse gases are set to zero (see Listing 1.10).

Listing 1.10: Radiation namelist radiation_nml for RCE in exp.atm_rce_test

```
&radiation_nml
irad_h2o
            = 1 ! prognostic vapor, liquid and ice
            = 2 ! constant co2
irad_co2
            = 0 ! no ch4
irad_ch4
irad_n2o
            = 0 ! no n2o
irad_o3
            = 4 ! perpetual january of ozone file linked
irad_o2
            = 0 ! no o2
irad_cfc11 = 0 ! no cfc11
irad_cfc12 = 0 ! no cfc12
irad_aero = 0 ! no aerosol
                       ! 4 if a diurnal cycle is switched on
isolrad
             = .FALSE. ! no diurnal cycle, .TRUE. for diurnal cycle
ldiur
```

1.6 Running the Model (Real Case)

The ICON code, as checkout from the SVN repository, does not include runscripts. Instead the run directory (\$ICONDIR/run/) includes several descriptor files for building grids, defining experiments and post-processings. There exist three different types of descriptor files with prefixes grid, exp, post:

- grid.<name>: to configure the grid generator, see chapter 1.8 for more details. It is recommended to use pre-built grids. For details, see section 1.7.
- exp.<name>: to define the namelist, which determinate the experiments.
- post.<name>: to define post-processing.

1.6.1 Input Data

Generally ICON requires the following input data: Grid files, external parameters, initialization (DWD analysis or IFS), input fields for radiation.

Grid Files

In order to run ICON, it is necessary to have the horizontal grid information as an input parameter. This information is stored within so-called grid files. For a ICON run, one global grid file is necessary. Additionally, if you want to nest, grid files of the nested domains are necessary, too. To improve the performance of ICON, a (optional) reduced radiation grid for each domain may be used.

The naming of the ICON-Grid is as follows: The initial icosahedron grid is refined by <n>-secting the edges, and further refinement is obtained by iteratively bisecting the created edges. The grid produced at the <k>refining iteration is named "R<n>B<k>". For further details, see the ICON Technical Documentation.

It is recommended to use pre-built grids. Further information can be found in chapter 1.7. For building own grids, the reader is referred to chapter 1.8. The names of the grid files have to be specified within the grid_nml:

```
&grid_nml
dynamics_grid_filename = "<INSERTFILENAME>"
radiation_grid_filename = "<INSERTFILENAME>"
```

External Parameters

ICON requires geographical localized datasets like the topographic height of the earth surface, the plant cover, the distribution of land and sea and, dependent on the schemes used, a variety of other so called external parameters. The EXTPAR software system (EXTPAR - External Parameter for Numerical Weather Prediction and Climate Application) is able to generate external parameters for the different models GME, COSMO, HRM and ICON. The software can run on a UNIX or Linux systems where the raw data is stored. It allows operators (experienced users) running the scripts to create new external parameters controlled by user specifications like the model domain. For a more detailed overview of EXTPAR, the reader is referred to the User and Implementation Guide of EXTPAR.

The name of the EXTPAR file which has to be read by ICON can be specified as follows:

```
extpar_filename = "<INSERTFILENAME>"

If not specified explicitly, ICON uses the following file name:
"<path>extpar_<gridfile>".
<path> and <gridfile> are then replaced at runtime by ICON.
```

&extpar_nml

Initialization

For the initialization of ICON, input data from either DWD or IFS is needed.

In case of DWD (init_mode=1) a first guess and an analysis is required:

```
&initicon_nml
dwdfg_filename = "<INSERTFILENAME>"
dwdana_filename = "<INSERTFILENAME>"
```

If not specified explicitly, ICON uses the following file names:

```
"<path>dwdFG_R<n>B<k>_DOM<idom>.nc" and
```

<path>, <n>, <k> and <idom> are then replaced at runtime by ICON according to the chosen
gridfile (see 1.6.1). The variable <idom> is an index for the domain on which the calculations
are performed. <idom>=0000 is reserved for a reduced radiation grid, <idom>=0001 for the global
domain, higher numbers are used for nested domains. NETCDF as well as GRIB2 input can be
used.

In case of IFS (init_mode=2) an analysis is required. It has to be in NetCDF:

```
&initicon_nml
ifs2icon_filename = "<INSERTFILENAME>"
```

If not specified explicitly, ICON uses the following file name:

```
"<path>ifs2icon_R<n>B<k>_DOM<idom>.nc".
```

<path>, <n>, <k> and <idom> are then replaced at runtime by ICON according to the chosen
gridfile (see 1.6.1). The variable <idom> is an index for the domain on which the calculations
are performed. <idom>=0000 is reserved for a reduced radiation grid, <idom>=0001 for the global
domain, higher numbers are used for nested domains.

Radiation

ICON requires input fields for the RRTM radiation scheme. The file names are specified as follows:

```
&nwp_phy_nml
lrtm_filename = "<INSERTFILENAME>"
cldopt_filename = "<INSERTFILENAME>"
```

If not specified explicitly, ICON uses the following file names:

```
"rrtmg_lw.nc" and
```

The files can be found within \$ICONDIR/data.

1.6.2 Creating a Runscript

To create a runscript, new users are advised to use the namelist descriptor file exp.nh-oper which contains recently recommended namelist settings. It might be necessary to account for the file

[&]quot;<path>dwdana_R<n>B<k>_DOM<idom>.nc".

[&]quot;ECHAM6_CldOptProps.nc".

names and paths of the input data. Additionally, machine dependent settings need to be added to this script to obtain a runscript. For some architectures, this step can be performed by using the make runscript environment as shown in 1.6.4. In the following, example settings for DWD CRAY are listed.

```
CRAY EXAMPLE: Namelists
<<Place your namelists e.g. from exp.nh_oper here>>
```

```
CRAY EXAMPLE: Submitting a job
aprun

-n <<INSERT: Total number of MPI Tasks>> \
-N <<INSERT: MPI Tasks/Node>> \
<<INSERT: Hyperthreading e.g. 2 -> 20 physical -> 40 "virtual" cores>> \
-d <<INSERT: Threads/MPI Task>> \
-m <<INSERT: Amount of memory to use>> control_model
```

1.6.3 Restart

A restart of the model requires a restart file that has to be created by a previous model run. In the following the procedures and the corresponding namelist settings are explained.

Creating the initial restart file:

The first job in a series of model runs creates the first restart file. To do so we have to use the following namelist switches.

```
&master_nml
lrestart = .FALSE.
```

In addition we have to prescribe at which time interval the job should produce a restart file:

```
&io_nml
dt_checkpoint = "<Insert time in seconds>"
```

The ICON run then creates restart files for each domain 1, ..., n_dom, and for each restart output time step.

The filenames are generic and look like:

```
"<gridfile>_restart_<modeltype>_<timestamp>.nc",
```

An example would be:

```
"iconR2B06_D0M01_restart_atm_20110101T001200Z.nc" (NetCDF format)
```

This filename can be customized using the namelist parameter:

```
&mo_run_nml
restart_filename = "<INSERTFILENAME>"
```

This file contains:

- data
- namelists
- several attributes

Note: - ICON reads the namelists only once and assumes that these are identical for all domains. - Since we do not know about the total number of domains at startup, we have to ask the current restart file for the attribute "n_dom".

For each domain 1, ..., n_dom, a symbolic link is generated with the generic name:

```
"restart_<modeltype>_DOMxx.nc"
```

Note: - The domain-dependent suffix "...DOMxx" is also required for non-nested setups.

Running the model in the restart mode:

ICON has to be informed that you want to carry out a restart run:

```
&master_nml
lrestart = .TRUE.
```

The generic link "restart_<modeltype>_DOMxx.nc" is used by the restart run to point to the last written restart file of the previous model run.

Chain of restart runs

If a chain of restart runs is foreseen it is recommended to use the namelist parameter dt_restart.

```
&time_nml
dt_restart = "<Insert time in seconds>"
```

In this case only one restart file is produced by each model run and after writing the restart file the job stops.

Note:- dt_restart and dt_checkpoint have to be selected carefully.

Asynchronous restart:

The restart can be handled by separated processors. The number of restart processors can be chosen by the user. The corresponding namelist parameter is:

```
&parallel_nml
num_restart_procs = n
```

n is the number of processors used for restart.

1.6.4 Make Runscript Environment

A full listing of descriptor files you will find in \$ICON/run/.

After configuration and compiling (chapter 1.3) these descriptor files can be transformed into runscripts, which should include the necessary system dependent parameters and the execution section exec.icon (\$ICONDIR/run/exec.iconrun), which starts the actual integration. This transformation is done in \$ICONDIR by:

```
./make_runscripts
```

This transforms every existing descriptor file in \$ICONDIR/run/<type>.<name> into a ready-to-use run script \$ICONDIR/run/<type>.<name>.run

For illustration there exists also

```
./make_my_runscripts
```

which transforms a single descriptor file into a run script. This file is an exemplary file and you can see how to define run parameters.

An exemplary descriptor file for a operational run is exp.nh_oper.

Note: if you change, or create a descriptor you will need to (re)create the run script in order for the changes to take effect.

To run a script <type>.<name>.run, either for creating grids or making an experiment or doing post-processing, go to the ./run folder

cd run

and use the job submission command, which depends on your machine:

```
[<submit>] <type>.<name>.run
```

[<submit>] is something like: {llsubmit,qsub}

Note: <u>Before</u> (!) running an experiment, the ICON grids must be available to the model. For this purpose, either pre-built grids and ExtPar Data can be used (see Sec. 1.7) or create own grids (1.8). For a new user, it is suggested to use pre-built grids first.

1.7 Pre-built Grids and ExtPar Data

A list of grid files has been pre-built for the ICON model together with the corresponding reduced radiation grids and the external parameters.

1. The **primary storage** location for ICON grids is

```
blizzard:/pool/data/ICON/grids/public
```

- 2. Every 24h the contents of the primary storage directory are mirrored to DWD's HPC.
- 3. Every 24h the contents of the primary storage directory are mirrored to a public web site: http://icon-downloads.zmaw.de.

Each grid file consists of a NetCDF file and a GPG signature file (http://de.wikipedia.org/wiki/GNU_Privacy_Guard).

The signature file makes sure that a grid file is complete and verifies the authorship.

1.7.1 Grid file nomenclature

The grids are identified by

- a **centre** number
- a **subcentre** number
- a numberOfGridUsed which is simply an integer number, increased by one with every new 'official' grid.

The grid files and the external parameter files are named accordingly, e.g.,

```
icon_grid_0001_RxxByy_G.nc
icon_extpar_0001_RxxByy_G.nc
```

where the name components are as follows:

The numberOfGridUsed parameter is part of the file name (0001, ...) and makes this file name unique.

In general, a lookup table is required to find the actual file name to which a set of these parameters corresponds. This 'table file' is located under

```
http://icon-downloads.zmaw.de/dwd_grids.xml
```

(the table file itself is under version control: https://svn.zmaw.de/svn/icon_grid_table).

1.8 Grid Generation

1.8.1 ICON atmosphere grids

The ICON horizontal spherical grid is based on the projection of the icosahedron on the sphere. This is a 2-dimensional grid, representing the earth's surface. The ICON grids need to be created, stored as NetCDF files, and consequently used by the ICON model. Alternatively, already stored grids may be used.

The initial icosahedron grid is refined by <n>-secting the edges, and further refinement is obtained by iteratively bisecting the created edges. The grid produced at the <k>refining iteration is named "R<n>B<k>", and the corresponding NetCDF-file is "iconR<n>B<k>-grid.nc". The grid files, after their creation, are located in the ./grids folder. For more detailed information about horizontal ICON grids the reader is referred to the ICON technical documentation.

Examples of grids are in ./grids. More information can be found in: \$ICONDIR/doc/technical/icon_grid.pdf.

The example given below shows the namelist parameters for generating a global R2B6 grid.

```
EXAMPLE Grid Generation of a R2B6 grid

#!/bin/ksh

#------

# Creation of atmosphere grids for ICON

#------

# ICON grid generator namelist parameters

#

# For a complete list see Namelist_overview and Namelist_overview.pdf

#

# nroot = Number of sections into which the edges of the original icosahedron are divided in the initial refinement step.
```

```
#
                (icosahedron = "grid -1" --> "nroot" grid = grid "0")
#
# grid_levels = Number of refinement steps applying edge bisection,
               follows the initial "nroot" refinement step.
#
                (grid "0" --> grid "1" --> ... --> grid "grid_levels")
#
# itype_optimize grid optimization method applied from grid level 1 onward.
#
               i | optimization | suffix fo grid output file
#
                _____
               0 | none
                                  l noo
                1 | Heikes Randall | hro
#
#
                4 | spring dynamics | spr
# beta_spring = Tuning parameter for spring dynamics to be chosen in the
#
              range [0.9,1.1]. Weights the target length between the
#
              grid points.
#
#-----
# First generate graphs
R=2
      # nroot (the first dissection will be a bisection)
      # highest grid level to reach (number of consequent bisections)
maxlev_optim=6 # highest grid level to apply optimizations
cat > NAMELIST_GRAPH << EOF
&graph_ini
            = \$\{R\}
 nroot
 grid_levels = ${B}
EOF
echo global_graph_generator null > $commandFile
${start} ${run_commmand}
check_error $? "global_graph_generator"
#-----
# Generate grids using the spring dynamics optimization
cat > NAMELIST_GRID << EOF
&grid_ini
            = \$\{R\}
 grid_levels = ${B}
&grid_options
 itype_optimize = 4    ! 1 = Heikes-Randall, 4 = spring dynamics
 maxlev_optim = $maxlev_optim ! the maximum level to optimize
EOF
```

ICON gives the possibility to nest subdomain within a parent grid. The example below gives the namelist parameters for generating a nested grid (patch). The root bisection of the patch in this

example starts with the fourth level of the bisections of the global model.

```
EXAMPLE Nested grid based on the global grid described before.
# Next the patches will be created.
# If the pathes are not needed then uncomment the next exit command
#exit
#-----
# ICON prepare_gridref namelist
# Parameter overview:
# grid_root: Number of root bisections
# start_lev: Grid level of global domain
#
# n_dom:
           Total number of model domains (including the global one)
# parent_id: List of parent domain ID's (starts at first nested domain,
            which has ID=2)
           true = circular subdomains, false =
# l_circ:
           rectangular (lat/lon) subdomains
#
# l_rotate: true: rotate center point into equator in case of l_circ=false
            this yields truly rectangular subdomains, whereas subdomains
            are conical otherwise because of the convergence of meridians
#
# NOTE:
           For subdomains crossing a pole, either l_circ=true
           or l_rotate=true is required
# l_plot:
           true: Generates GMT files for domain configuration
# NOTE:
            The following parameters have to be specified for each nested
            domain!
# radius:
           radius (deg) of nested domains (for l_circ=true)
# center_lon: Center longitude of nested domains
# center_lon: Center latitude of nested domains
# hwidth_lon: half-width longitude of nested domains (for l_circ=false)
# hwidth_lat: half-width latitude of nested domains (for l_circ=false)
#
  _____
```

```
# suffix of grid files which specifies optimization type
# (without optimization leave empty)
#OPTFIX=spr0.90_M4
# NOTE: _M4 means that maxlev_optim = 4 has to be set in the grid generator
# maxlev_optim is not needed for Heikes-Randall optimization
# Create plots of domain configuration
PLOTS=.false.
cat > NAMELIST_GRIDREF << EOF
&gridref_ini
 grid_root = 2
 start_lev = 4
          = 2
 n\_dom
 parent_id = 1,
 1_circ
           = .true.
            = .true.
 l_rotate
 1_plot
            = .true.
            = 30.,
 radius
 center_lon = -90.,
 center_lat = 40.,
 hwidth_lon = 55.,
 hwidth_lat = 55.,
 bdy_indexing_depth = 14
EOF
```

1.8.2 Information contained in grid files

The ICON grids are treated as a general unstructured grid, so the grid NetCDF-files contain the full information of the location and the connectivity of all the grid entities (cells, edges and vertices). The grid nesting hierarchy information is also included.

Some basic variables that may be useful for plotting are:

```
double clon(cell)
                                : longitude of cell centers [radian]
double clat(cell)
                                : latitude of cell centers [radian]
double clon_vertices(cell, nv) : longitudes of the vertices of the cell [radian]
double clat_vertices(cell, nv) : latitudess of the vertices of the cell [radian]
double elon(edge)
                                : longitude of edge midpoint [radian]
double elat(edge)
                                : latitude of edge midpoint [radian]
double elon_vertices(edge, no) : longitudes of the vertices of the edges [radian]
double elat_vertices(edge, no) : latitudes of the vertices of the edges [radian]
double vlon(vertex)
                                : longitude of vertices [radian]
double vlat(vertex)
                                : latitude of vertices [radian]
                                : area of grid cell [m2]
double cell_area(cell)
double cell_elevation(cell) : elevation at the cell centers [m]
      cell_sea_land_mask(cell): sea (-2 inner, -1 boundary)
```

```
land (2 inner, 1 boundary) mask for the cell
...

double edge_length(edge) : lengths of edges of triangular cells [m]
double dual_edge_length(edge) : lengths of dual edges (distances between triangular cell circumcenters) [m]
...
```

For a full listing of variables contained in a grid file, for instance in iconR2B04-grid.nc, use:

```
ncdump -h iconR2B04-grid.nc
```

or

cdo sinfov iconR2B04-grid.nc

1.8.3 Viewing/plotting grids

In order to plot an icon grid you should ensure that ncl-6.0 and cdo-1.5.4 is available on your machine. Then go to the \$ICONDIR/grids/ folder and give:

```
alias iplot="ncl $ICONDIR/scripts/postprocessing/tools/icon_plot.ncl
'altLibDir="$ICONDIR/scripts/postprocessing/tools/"'" iplot 'iFile="<grid file name>"'
'mapType="ortho"' 'varName="cell_sea_land_mask"' 'oType="png"' 'showGrid=True'
'lStrg="Cell sea land mask"' 'bStrg=""'
```

The above example will plot cell sea land mask. More details on plotting can be found at the Visualization chapter.

The \$ICONDIR/run/post.plot_icon_grids script can be used to plot nested grids. Go to \$ICONDIR/run/folder and give:

```
./post.plot_icon_grids
```

A PDF-file with a plot of the iconR2B04_DOM01 and iconR2B05_DOM02 grids will appear on your screen. (Note that this process is time consuming.)

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2 Output

In general the user has to specify six individual quantities to generate output of the model. These are:

- 1. The time interval between two model outputs.
- 2. The name of the output file.
- 3. The name of the variable.
- 4. The type of the vertical output grid (e.g. pressure levels or model levels).
- 5. The type of the horizontal output grid (e.g. ICON grid or geographical coordinates).

ICON offers the possibility to write groups of variables. In the following we will present two examples to demonstrate the options the user has to prescribe these quantities. A detailed description of all namelist parameters available to organize the output is described in io_nml in the namelist section.

Example 1

We will begin with an individual variable which is written in NETCDF format on pressure levels and is interpolated to a horizontally regular lat-long grid:

```
NAMELIST EXAMPLE
&io_nml
filetype
                                   ! output format: 2=GRIB2, 4=NETCDFv2
dom
                                  ! write output for domain 1
                              0., 1.E7, 3600. ! start, end, interval in s.
output_bounds
                                  ! max. num. of time steps within one file
steps_per_file
mode
                                  ! 1: forecast mode (relative t-axis)
include_last
                           = .TRUE. ! include the last time step
output_filename
                           = '<INSERTFILENAME>' ! file name base
pl_varlist
                           = 'geopot' ! name of pressure level field
                                  ! output is transferred to lat long grid
remap
reg_lon_def
                           = 0.,0.5,359.5
                                             !start, incr., end, in deg.
                           = 90.,-0.5, -90. !start, incr., end, in deg.
reg_lat_def
```

Example 2

The flexibility of the options ICON offers is demonstrated in another example. Now we apply an alternative to define the runtime of ICON, write several variables, at the same time, in one data

set, on model levels, and on the original horizontal grid of ICON. In addition the example below shows the options when several model domains run at the same time and we want to produce output for all model domains.

```
NAMELIST EXAMPLE
&output_nml
dom
                                 -1 ! write all domains
steps_per_file
                                 5 ! max. num. of time steps within
  output_start
                   = "1978-01-01T00:00:00Z" ! ISO-format date+time
                   = "1979-01-02T00:00:00Z" ! ISO-format date+time
  output_end
  output_interval
                   = "PTO1H"
                                             ! ISO-format interval
  file_interval
                   = "PTO1D"
                                             ! ISO-format interval
  include_last
                   = .FALSE.
  output_filename
                               = '<INSERTFILENAME>'
                                                        ! file name base
 ml_varlist='u', 'group:precip_vars' ! Indiv. variable and variable group
                   = .TRUE. ! Output on the ICON horizontal grid
  output_grid
```

Variable groups

Next we explain the meaning of variable groups. Using the "group:" keyword for the namelist parameters ml_varlist, hl_varlist, pl_varlist, sets of common variables can be added to the output.

There exists a special syntax which allows to remove variables from the output list, e.g. if these undesired variables were contained in a previously selected group.

Typing "-<varname>" (for example "-temp") removes the variable from the union set of group variables and other selected variables. Note that typos are not detected but that the corresponding variable is simply not removed!

How to find variable names and contents of variable groups

Finding the correct names of the variables you may want to write to a data set is not an easy task and you should be aware of some pitfalls. We will help you to avoid the most obvious ones. First of all users that have already experience with the COSMO model should know that the names of the atmospheric variables in ICON are **not identical**.

The easiest way to identify the correct names of the variables you would like to write is to look into the following data sets:

```
atm_dyn_iconam/mo_nonhydro_state.f90
atm_phy_nwp/mo_nwp_phy_state.f90
lnd_phy_nwp/mo_nwp_lnd_state.f90
```

Now you may want to use the option of writing groups of variables and of course you may want to know which variable belongs to which group. Keep in mind that there is an option mentioned before to remove variables from the output of a group of variables.

The following table gives overview on the allocation of variables to individual variable groups. If

you want to translate the Fortran variables to the physical or mathematical ones again have a look to the Fortran files listed above.

```
*********
       nh_prog_vars
vn
rho
theta_v
exner
*********
       dwd_fg_atm_vars
vn
W
rho
theta_v
tke
u
pres_sfc
temp
pres
z_ifc
t_2m
td_2m
u_10m
v_10m
*********
       mode_dwd_fg_in
vn
W
rho
theta_v
tke
t_g
t_mnw_lk
t_wml_lk
h_ml_lk
t_bot_lk
c_t_{lk}
t_b1_lk
h_b1_lk
qv_s
w_i
w_so_ice
w_snow
rho_snow
t_snow_mult
rho_snow_mult
wliq_snow
wtot_snow
```

```
dzh_snow
gz0
*********
     atmo_ml_vars
W
tke
u
temp
pres
*********
     atmo_pl_vars
tke
u
temp
*********
     atmo_z1_vars
W
tke
u
temp
pres
*********
     mode_dwd_ana_in
u
٧
temp
pres
t_ice
h_ice
fr_seaice
w_so
t_snow
h_snow
freshsnow
*********
     atmo_derived_vars
omega
div
vor
*********
     land_vars
t_g
qv_s
w_i
w_p
```

```
w_s
t_so
w_so
w_so_ice
t_snow
w_snow
rho_snow
snowfrac
*********
       dwd_fg_sfc_vars
t_g
t_ice
h_ice
fr_seaice
w_i
t_so
w_so
w_so_ice
t_snow
w_snow
rho_snow
h_snow
freshsnow
t_snow_mult
rho_snow_mult
wliq_snow
wtot_snow
dzh_snow
gz0
*********
       mode_combined_in
t_g
t\_ice
h_ice
qv_s
fr_seaice
w_i
w_so
t_snow
w_snow
rho_snow
h_snow
freshsnow
*********
       mode_cosmode_in
t_g
t_ice
h_ice
qv_s
```

```
w_i
w_so
t_snow
w_snow
rho_snow
h_snow
freshsnow
*********
       dwd_fg_scf_vars
t_mnw_lk
t_{wml_lk}
h_ml_lk
t_bot_lk
c_t_{lk}
t_b1_lk
h_b1_lk
qv_s
********
       land_tile_vars
t_g_t
t_s_t
w_i_t
w_p_t
w_s_t
t_so_t
w_so_t
w_so_ice_t
t_snow_t
w_snow_t
rho_snow_t
t_snow_mult_t
wtot_snow_t
wliq_snow_t
rho_snow_mult_t
dzh_snow_t
qv_s_t
h_snow_t
snowfrac_t
snowfrac_lc_t
*********
       snow_vars
t_snow
rho_snow
wliq_snow
wtot_snow
dzh_snow
*********
       multisnow_vars
t_snow_mult
```

```
rho_snow_mult
wliq_snow
wtot_snow
dzh_snow
*********
      precip_vars
rain\_gsp
snow_gsp
rain_con
snow_con
ice_gsp
graupel_gsp
hail_gsp
tot_prec
*********
      additional_precip_vars
con_prec_rate_avg
gsp_prec_rate_avg
cape
clct
tot_cld_vi
*********
      pbl_vars
gust
shfl_s
lhfl_s
lhfl_bs
lhfl_pl
ghfl_s
tcm
tch
{\tt t\_2m}
qv_2m
td_2m
u_10m
v_10m
tkvm
tkvh
*********
      cloud_diag
clc
gc_dia
gi_dia
tot_cld
**********
      rad_vars
thb_s
sod_t
```

```
sou_t
sod_s
sou_s
thd_s
thu_s
sodird_s
sodifd_s
sodufu_s
albdif
albvisdiff
albnirdiff
sob_s_t
thb_s_t
flxdwswtoa
sob_s
sob_t
*********
       phys_tendencies
ddt_temp_radsw
ddt_temp_radlw
ddt_temp_turb
ddt_temp_drag
ddt_u_turb
ddt_u_so
ddt_u_gwd
ddt_v_turb
ddt_v_sso
ddt_v_gwd:
*********
       prog_timemean
temp_m
rho_m
u_m
v_m
pres_sfc_m
pres_msl_m
*********
       echam_timemean
cosmu0_m
flxdwswtoa_m
aclcov_m
rsfl_m
rsfc_m
ssfl_m
ssfc_m
totprec_m
qvi_m
xlvi_m
xivi_m
```

```
swflxsfc_m
swflxtoa_m
lwflxsfc_m
lwflxtoa_m
tsfc_m
evap_m
lhflx_m
shflx_m
u_stress_m
v_stress_m
**********
       tracer_timemean
qc_m
qv_m
qi_m
*********
       atmo_timemean
all vars of prog_timemean, echam_timmean, tracer_timemean
```

Data format

ICON offers the possibility to produce output either in NETCDF or GRIB2 format. This can be chosen by the namelist parameter filetype of the namelist &output_nml. New users are suggested to set filetype=4 in order to use NETCDF output.

In GRIB2, a variable is uniquely defined by the following set of metadata:

- Discipline (see GRIB2 code table 4.2)
- ParameterCategory (see GRIB2 code table 4.2)
- ParameterNumber (see GRIB2 code table 4.2)
- typeOfFirstfixedSurface and typeOfSecondFixedSurface (see GRIB2 code table 4.5)
- step Type (instant, accum, avg, max, min, diff, rms, sd, cov, ...)

Time stamp format

The namelist parameters output_start, output_end, output_interval allow the specification of time stamps according to ISO 8601. The general format for time stamps is YYYY-MM-DDThh:mm:ss where Y: year, M: month, D: day for dates, and hh: hour, mm: minute, ss: second for time strings. The general format for durations is PnYnMnDTnHnMnS. See, for example, http://en.wikipedia.org/wiki/ISO_8601 for details and further specifications.

NOTE: as the mtime library underlaying the output driver currently has some restrictions concerning the specification of durations:

- 1. Any number n in PnYnMnDTnHnMnS must have two digits. For instance use "PT06H" instead of "PT6H"
- 2. In a duration string PnyearYnmonMndayDTnhrHnminMnsecS the numbers nxyz must not pass the carry over number to the next larger time unit: 0<=nmon<=12, 0<=nhr<=23, 0<=nmin<=59, 0<=nsec<=59.999. For instance use "PT01D" instead of "PT24H", or "PT01M" instead of "PT60S".

Soon the formatting problem will be resolved and the valid number ranges will be enlarged. (2013-12-16).

Extra output

- 1. In the namelist run_ctl set the number of fields with inextra_2d or inextra_3d. The logical variable for output lwrite_extra then will be set automatically. Note, the number of extra fields is limited by 9 each for 2D and 3D.
- 2. USE these variables in the module needed.
- 3. Implement the storage of wished fields by using the nonhydrostatic diagnostic type with p_diag%extra_2d/3d.

Example for the use of p_diag%extra_2d:

```
USE mo_global_variables, ONLY: inextra_2d
...
DO jc = i_startidx, i_endidx
p_diag\%extra_2d(jc,jb,1)= yxz(jc,jb)
ENDDO
```

Asynchronous output:

It is highly recommended that the asynchronous output option of ICON is applied. In short this option reserves a number of processors for output only. While writing the remaining processors continuously carry out calculations. Otherwise they would have to wait until output is finished. The corresponding namelist parameter is:

```
&parallel_nml
num_io_procs = n
```

n is the number of processors used for output.

Time mean output:

The builtin functionality for getting time-averaged output fields has the following features and limitations

- The list of variables is configurabe via output_nml like the regular instantaneous
- There can be multiple mean values intervals for the same variable, but only one interval per file
- 2d and 3d Variables are supported
- output is limited to model level
- nesting is not supported
- GRIB output is not supported
- Output interval has to be a divisor of the restart interval, because the intermediate accumulation results are not saved
- The output for the initial timestep of the time mean variables is zero.

Example Usage For getting 6-hourly averaged output, two namelist variable of output_nml has to be set up:

```
• output_interval = "PTO6H"
```

• operation = "mean"

If daily mean values should be created in addition, a new output_nml has to be defined for a new output file with output_interval = "P1D".

Discussion

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3 Visualization

Visualizing data on a non-regular grids is a task on its own, because the number of tools for solving such problem is very limited. NCL is one of them and we chose it as the main tool for ICON. You can find several examples of how to write simple plot scripts for ICON data sets on this website: http://www.ncl.ucar.edu/Applications/icon.shtml. The coordinate information is essential for writing your own plot scripts. ICON output files currently have three different types of them: cells, edges and vertices, e.g. tracers like temperature and salinity and surface elevation are defined on each cell center while the normal velocity is defined on edges.

3.1 icon_plot.ncl

For getting around the different coordinates and in order not to rewrite things there is a general plot scripts: icon_plot.ncl. It supports contour and vector plots, a combination of both via overlaying and vertical sections. Both atmosphere and ocean vertical coordinate systems can be handled by it: While ocean uses a plain depth axes, atmosphere model uses hybrid sigma pressure levels (hydrostatic) and free 3D height variable (non-hydrostatic).

The script icon_plot.ncl is a single NCL program, which provides multiple plot types for data on ICON's grid. It is located in the ICON-repository under

source:/trunk/icon-dev/scripts/postprocessing/tools/icon_plot.ncl. Most of the functionality is implemented in a library: icon_plot_lib.ncl located in

source:/trunk/icon-dev/scripts/postprocessing/tools/icon_plot_lib.ncl. Both files are installed into the /pool/data/ICON/tools which is the default lookup location for the library. For different location like an icon checkout, use altLibDir, e.g.

altLibDir='"/home/user/src/icon-dev/tools"'.

3.1.1 Requirements

- NCL 5.2.1 is the minimum version of NCAR's plotting language (http://www.ncl.ucar.edu)
- CDO (https://code.zmaw.de/projects/cdo)

3.1.2 Customization

icon_plot.ncl optionally reads a configuration file named \$HOME/.icon_plot.rc where default options can be set. Actually it is handled like an ordinary ncl file. This can be used to customize the altLibDir setting, e.g.:

altLibDir="/home/ram/src/git/icon/scripts/postprocessing/tools"
oType="png"

3.1.3 Basic command line option

Required are options for

1. Input/output files: Use the variable iFile for defining the input and oFile for the output file. It's extension depends on the output type, which can be set with oType. If oFile is left out, the output file will inherit its name from the input file.

2. Variable selection: Depending on the plot mode you like to use, varName for scalar variables or vecVars for vector-variables must be uses.

Optional (default:0) parameter are

- 1. **Level selection**: Levels can only be selected by their index. That's why, the corresponding variables is called **levIndex**. Please note that it starts with 0, like any other NCL indices.
- 2. **Time selection**: Like levIndex, the variable timeStep can be used to select a certain time step, again starting from 0.

There are many more parameters (see 3.1.8) for mapping, transections, selecting regions and masking, but these are the most fundamental ones.

3.1.4 Plot Types

For flexibility the selection of a specific plot mode is implemented by combining certain options.

Contour plots

Contour plot are the default plot mode. If only the require parameters are set, e.g. iFile and varName, a simple contour plot is created with

```
ncl icon_plot.ncl 'varName="T"' 'iFile="iFILENAME"'
```

This is a basic temperature plot. Captions are set to basic information like variable name, time and level information and input filename.

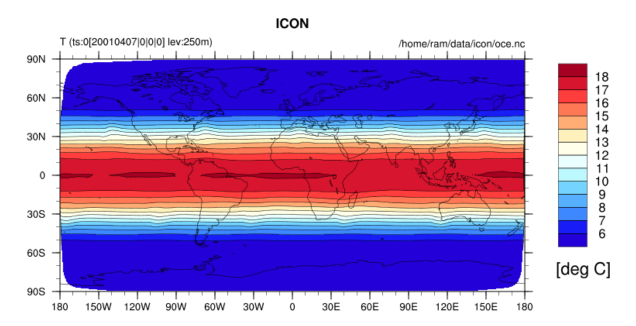
Vector plots

Use vecVars instead of varName. To adjust the length of the reference vector, use the variable vecRefLength.

```
ncl icon_plot.ncl 'vecVars="U V"' 'iFile="iFILENAME"' vecRefLength=0.01
```

Overlay of scalar and vector variables

Contour and vector plots can be combined into a single plot by overlaying both. Following this approach, such an overlay plot will be created, if varName and vecVars are given:



Prgr icon_plot.ncl: Wed Nov 30 14:28:49 CET 2011,ram

Figure 3.1: Example of contour plot

ncl icon_plot.ncl 'varName="T"' 'iFile="iFILENAME"' 'vecVars="U V"'

Vertical sections

Data for sections have to be interpolated first. This is done internally and you do not have to care about it. Section plot are created, if a start and and end point of a section is given. For this purpose, the variables secLC (section-left-corner) and secRC (section-right-corner) have to be used. Theses variable have to be (lon,lat) arrays like secLC=(/20.,30./).

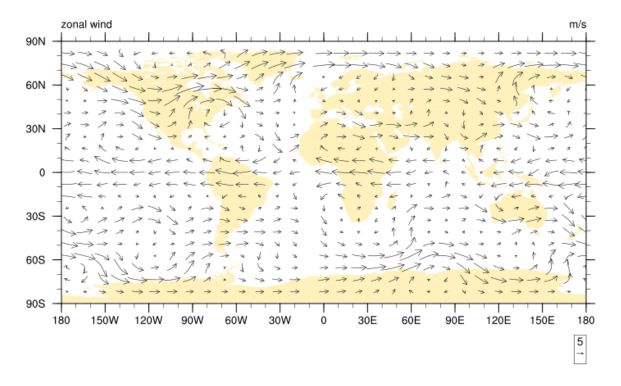
Example call:

```
ncl icon_plot.ncl 'varName="T"' 'iFile="iFILENAME"' \
    'secLC=(/0,80/)' 'secRC=(/0,-80/)'
```

secPoints is an option to set the accuracy of the plot. The representing of the location of the section is suppressed by setting showSecMap=False. Its default value is True.

Display the ICON grid

Set the parameter **showGrid** to **True** and for scalar variables, the ICON grid is represented instead of the contour plot. For large grids, this can take a long time.



Prgr icon_plot.ncl: Wed Nov 30 14:39:10 CET 2011,ram

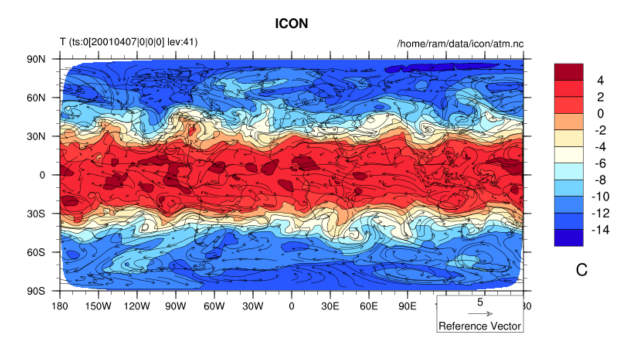
Figure 3.2: Example of vector plot

3.1.5 Regional plots

Use the variables mapLLC (map-Lower-Left-Corner) and mapURC (map-Upper-Right-Corner) to select special regions of the earth. Here is a list of useful examples:

Table 3.1: Examples of useful regional plots

Trop. Atlantic	'mapLLC=(/-60, -25/)'	'mapURC=(/ 25,25/)'
North Polar	'mapLLC=(/-200, 20/)'	'mapURC=(/160,90/)'
North Atlantic	'mapLLC=(/-100,-15/)'	'mapURC=(/ 35,65/)'
Labrador/Panama	'mapLLC=(/-200, -5/)'	'mapURC=(/ 35,85/)'
North Atlantic/Eurasia	'mapLLC=(/ -80, -5/)'	'mapURC=(/ 75,85/)'
Asia	'mapLLC=(/ 20,-15/)'	'mapURC=(/160,85/)'



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Figure 3.3: Example of overlay plot

3.1.6 Masking

Masking can be done in two different ways:

1. Manually mask the data with CDO before running the plot scripts, i.e. use the ifthen operator or perform a division with the mask variable:

cdo div iconInput.nc -selname,mask_variable iconInput.nc maskedOutput.nc

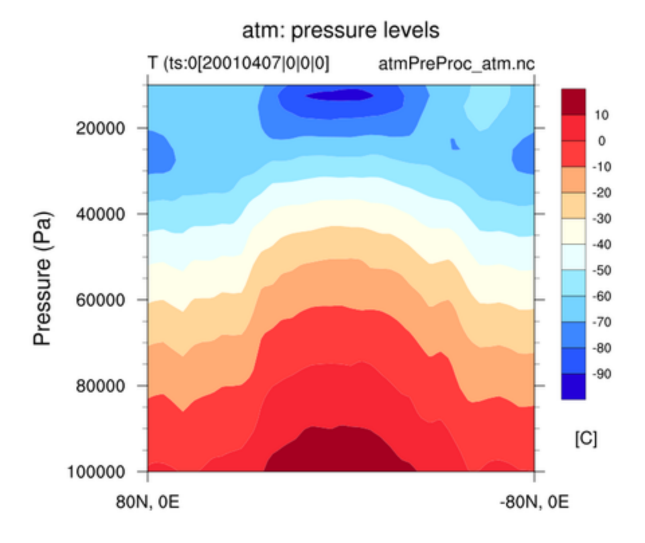
2. Let the plot script perform the masking using the NCL's mask function. For this purpose, the commandline variables maskName and maskFile have to be used. If the mask variable is part of the regular input file, maskFile can be left out.

Both methods have their pros and cons. Whereas the second methods works fine for all types of horizontal representation, the first produces better results for vertical cross sections.

3.1.7 Data on other grids

Although icon_plot.ncl is implemented for ICON, it can be used for data an regular grids, too. In this case, internal interpolation is not performed.

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Figure 3.4: Example of vertical sections plot

3.1.8 All options

 ${\tt icon_plot.ncl}$ has built-in documentation of all options. Use

ncl icon_plot.ncl help=True

3.2 ncview/GrADS

Neview (http://meteora.ucsd.edu/~pierce/neview_home_page.html) and GrADS (http://www.iges.org/grads/) can be used after converting icon data sets to a regular grid. This can easily be done with cdo:

cdo -P 8 -r remapnn,r180x90 icon.nc regular_icon.nc

This uses nearest neighbor interpolation and hereby keeps the model values. When using a higher regular resolution the triangular icon grid keeps visible.

3.3 Other Possibilities

- GMT is useful, when the grid should be visualized.
- ParaView is an alternative to display data on an unstructured grid. As a caveat, the model output has first to be converted into the vtk format.

Discussion

Document last edited by I. Kraut on 29.11.2013. Note: -

4 ICON Namelists Overview

4.1 Namelist Annotation

Every ICON run generates annotated lists of namelist parameters during the setup. These lists are written to text files nml.atmo.log, nml.cpl.log, nml.ocean.log and have the following form:

```
NAMELIST IO_NML
OUT_EXPNAME 'case4 [...]' (truncated)
>> DEFAULT: 'IIIEEEETTTT [...]' (truncated)
OUT_FILETYPE 2
LKEEP_IN_SYNC F
DT_DATA 43200.00000000000000000
DT_DIAG 1728000.00000000000
>> DEFAULT: 21600.00000000000
>> DEFAULT: 86400.000000000000
```

and so on.

The DEFAULT annotation denotes all those parameters that have been modified by the user; in this case the default value of the namelist parameter is stated together with the modified value. All other namelist parameters are listed only with their default values.

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4.2 ICON Namelists

4.2.1 Scripts, Namelist files and Programs

Run scripts starting the programs for the grid generation and the models are stored in run/. These scripts write namelist files containing the specified Fortran namelists. Programs are stored in < icon home>/build/<architecture>/bin/.

Table 4.1: Namelist files

Namelist file	Purpose	Made by script	Used by program
NAMELIST_GRAPH	Generate graphs	create_global_grids.run grid_command	grid_command
$NAMELIST_GRID$	Generate grids	create_global_grids.run grid_command	grid_command
NAMELIST_GRIDREF	Gen. nested domains	create_global_grids.run	grid_command
NAMELISTICON	Run ICON models	exp.jname;.run	control_model

4.2.2 Namelist parameters

The following subsections tabulate all available Fortran namelist parameters by name, type, default value, unit, description, and scope:

- Type refers to the type of the Fortran variable, in which the value is stored: I=INTEGER, L=LOGICAL, R=REAL, C=character string
- Default is the preset value, if defined, that is assigned to this parameter within the programs.
- Unit shows the unit of the control parameter, where applicable.
- Description explains in a few words the purpose of the parameter.
- Scope explains under which conditions the namelist parameter has any effect, if its scope is restricted to specific settings of other namelist parameters

Information on the file, where the namelist is defined and used, is given at the end of each table.

4.3 Namelist parameters for grid generation

4.3.1 Namelist parameters defining the atmosphere grid

graph_ini (NAMELIST_GRAPH)

Parameter	Type	Default	Unit	Default Unit Description	Scope
nroot	I	2		root subdivision of initial edges	
$\operatorname{grid_levels}$	I	4		number of edge bisections following the root	
				subdivision	
lplane	П	.FALSE.		switch for generating a double periodic planar	
				grid. The root level consists of 8 triangles.	

Defined and used in: src/grid_generator/mo_io_graph.f90

grid_ini (NAMELIST_GRID)

					ě
Parameter	Type	Default	Unit	Default Unit Description	Scope
nroot	I	2		root subdivision of initial edges	
grid_levels	Ι	4		number of edge bisections following the root	
				subdivision	
lplane	П	FALSE.		switch for generating planar grid. The root	
				level consists of 8 triangles.	
lread_graph	L	FALSE.		switch for reading graph information from	
				precomputed file; .TRUE. implies that the	
				graph generator needs to be executed in	
				advance	

Defined and used in: src/grid_generator/mo_grid_levels.f90

grid_options (NAMELIST_GRID)

Parameter	Type	Default	Unit	Description	Scope
x_rot_angle	\mathbf{R}	0.0	deg	Rotation of the icosahedron about the x-axis	
				(connecting the origin and $[0^{\circ}E, 0^{\circ}N]$)	
y_rot_angle	${ m R}$	0.0	deg	Rotation of the icosahedron about the y-axis	
				(connecting the origin and [90°E, 0°N), done	
				after the rotation about the x-axis.	
z_rot_angle	R	0.0	deg	rotation of the icosahedron about the z-axis	
				(connecting the origin and [0°E, 90°N), done	
				after the rotation about the y-axis.	
itype_optimize	I	4		Grid optimization type	
				0: no optimization	
				1: Heikes Randall	
				2: equal area	
				3: c-grid small circle	

Parameter	Type	Default	Unit	Default Unit Description	Scope
				4, 5: spring dynamics; 5 with optimization for	
				faster convergence	
l-c-grid	L	.FALSE.		C-grid constraint on last level	
$maxlev_optim$	Ι	100		Maximum grid level where the optimization is	i_type_optimize = 1 or
				applied	4
beta_spring	\mathbf{R}	0.90		tuning factor for target grid length	$i_{type_optimize} = 4$

Defined and used in: src/grid_generator/mo_grid_levels.f90

plane_options (NAMELIST_GRID)

Parameter	Type	Default	Unit	Description	Scope
tria_arc_km	${ m R}$	10.0	km	length of triangle edge on plane	lplane=.TRUE.

The number of grid points is generated by root level section and further bisections. The double periodic root level consists of 8 triangles. The spatial coordinates are $-1 \le x \le 1$, and $-\sqrt{3}/2 \le y \le \sqrt{3}/2$. Currently the planar option can only be used as an f-plane.

Defined and used in: src/grid_generator/mo_grid_levels.f90

gridref_ini (NAMELIST_GRIDREF)

Parameter	Type	De	Unit	fault Unit Description S	Scope
grid root	I	2		root subdivision of initial edges	
start_lev	П	4		number of edge bisections following the root	
				subdivision	
m-dom	П	2		number of logical model domains, including the	
				global one	
$ m n_phys_dom$	П	mop-u		number of physical model domains, may be	
				larger than n-dom (in this case, domain	
				merging is applied)	

Parameter	Type	Default	Unit	Description Scol	Scope
parent_id	I(n-phys-	i		ID of parent domain (first entry refers to first	
	dom-1			nested domain; needs to be specified only in	
				case of more than one nested domain per grid	
				level)	
logical_id	I(n-phys-	i+1		logical grid ID of domain (first entry refers to	
	dom-1			first nested domain; needs to be specified only	
				in case of domain merging, i.e. n_dom <	
				n-phys-dom)	
l_plot	Г	.FALSE.		produces GMT plots showing the locations of	
				the nested domains	
$_{ m l_circ}$	Г	FALSE.		Create circular (.T.) or rectangular (.F.)	
				refined domains	
l_rotate	ı	.FALSE.		Rotates center point into the equator in case of lcirc	lcirc=.FALSE.
				1-circ = .FALSE.	
write_hierarchy	I	1		0: Output only computational grids	
				1: Output in addition parent grid of global	
				model domain (required for computing physics	
				on a reduced grid)	
				2: Output all grids back to level 0 (required for	
				hierarchical search algorithms)	
lsep_gridref_info	Г	FALSE.		.TRUE.: write fields describing parent-child	
				connectivities into separate grid files	
uuid_sourcefile	C(n-dom)	'EMPTY'		If specified, provides the names of existing grid	
				files from which the unid shall be copied. If a	
				radiation grid is present, the first entry refers	
				to this grid.	
${ m bdy_indexing_depth}$	Ι	12		Number of cell rows along the lateral boundary	
				of a model domain for which the refin_ctrl fields	
				contain the distance from the lateral boundary;	
				needs to be enlarged when lateral boundary	
	_	_		nuaging is required for one-way nesung	

Parameter	Type	Default Unit	Unit	Description	Scope
radius	R(n_dom-	30.	deg	radius of nested domain (first entry refers to	lcirc=.TRUE.
	1)			first nested domain; needs to be specified for	
				each nested domain separately)	
hwidth_lon	$R(n_{-}dom-$	20.	deg	zonal half-width of refined domain (first entry	lcirc=.FALSE.
	1)			refers to first nested domain; needs to be	
				specified for each nested domain separately)	
hwidth_lat	$R(n_{-dom}$	20.	deg	meridional half-width of refined domain (first	lcirc=.FALSE.
	1)			entry refers to first nested domain; needs to be	
				specified for each nested domain separately)	
center_lon	$R(n_{-}dom-$	30.	deg	center longitude of refined domain (first entry	
	1)			refers to first nested domain; needs to be	
				specified for each nested domain separately)	
center_lat	$R(n_{-}dom_{-}$	90.	deg	center latitude of refined domain (first entry	
	1)			refers to first nested domain; needs to be	
				specified for each nested domain separately)	

Defined and used in: src/grid_generator/mo_gridrefinement.f90

gridref_metadata (NAMELIST_GRIDREF)

Parameter	Type	Default	Unit	Default Unit Description	Scope
number_of_grid_used	I(n_dom+	0		sets the number of grid used in the netcdf	
	1)			header; the number of entries must be	
				n_dom+1 since the first number refers to the	
				radiation grid	
centre	Ι	0		centre running the grid generator	
				78: EDZW (DWD)	
				252: MPIM	
subcentre	Ι	0		subcentre to be assigned by centre, usually 0	

Scope			
Description	Output name style	1: Standard: iconRXBXX_DOMXX.nc	2: DWD: icon_grid_XXXX_RXXBXX_X.nc
Unit			
Default	1		
Type	I		
Parameter	outname_style		

Defined and used in: src/grid_generator/mo_gridrefinement.f90

4.4 Namelist parameters defining the atmospheric model

Namelist parameters for the ICON models are organized in several thematic Fortran namelists controling the experiment, and the properties of dynamics, transport, physics etc.

4.4.1 coupling_mode_nml

Parameter	Type	Default	Unit	Description	Scope
coupled_mode	Γ	.FALSE.		.TRUE.: if yac coupling routines have to be	
				called	

Defined and used in: src/namelists/mo_coupling_nml.f90

4.4.2 diffusion_nml

Parameter	Type	Default	Unit	Description	Scope
lhdiff_temp	Г	.TRUE.		Diffusion on the temperature field	
lhdiff_vn	Γ	TRUE.		Diffusion on the horizontal wind field	
lhdiff_w	Γ	TRUE.		Diffusion on the vertical wind field	

Parameter	Type	Default	Unit	Description	Scope
hdiff_order	I	4 (hy-		Order of ∇ operator for diffusion:	Options 2, 24 and 42
		dro)		-1: no diffusion	are allowed only in the
		5 (NH)		2: ∇^2 diffusion	hydrostatic atm model
				3: Smagorinsky ∇^2 diffusion	(iequations $= 1$ or 2 in
				4: ∇^4 diffusion	dynamics_nml).
				5: Smagorinsky ∇^2 diffusion combined with ∇^4	
				background diffusion as specified via	
				hdiff_efdt_ratio	
				24 or 42: ∇ 2 diffusion from model top to a	
				certain level (cf. k2-pres-max and k2-klev-max	
				below); ∇^4 for the lower levels.	
lsmag-3d	L	.FALSE.		.TRUE.: Use 3D Smagorinsky formulation for	hdiff_order=3 or 5;
				computing the horizontal diffusion coefficient	itype_vn_diffu=1
				(recommended at mesh sizes finer than 1 km if	
				the LES turbulence scheme is not used)	
itype_vn_diffu	I	1		Reconstruction method used for Smagorinsky	iequations=3,
				diffusion:	hdiff_order=3 or 5
				1: u/v reconstruction at vertices only	
				2: u/v reconstruction at cells and vertices	
itype_t_diffu	I	2		Discretization of temperature diffusion:	iequations=3,
				$1: K_h \nabla^2 T$	hdiff_order=3 or 5
				$2: \nabla \cdot (K_h \nabla T)$	
k2_pres_max	$_{ m R}$	-99.	Pa	Pressure level above which ∇^2 diffusion is	$hdiff_{order} = 24 \text{ or } 42,$
				applied.	and dynam-
					$ics_nml:iequations = 1$
					or 2.
k2_klev_max	I	0		Index of the vertical level till which (from the	hdiff_order = 24 or 42 ,
				model top) ∇^2 diffusion is applied. If a positive	and dynam-
				value is specified for k2_pres_max, k2_klev_max	ics_nml:iequations = 1
				is reset accordingly during the initialization of	or 2.
		_		a model run.	

Parameter	Type	Default	Unit	Description	Scope
hdiff_efdt_ratio	\mathbf{R}	1.0		ratio of e-folding time to time step (or 2^* time	
		(hydro)		step when using a 3 time level time stepping	
		36.0		scheme) (for triangular NH model, values above	
		(NH)		30 are recommended when using hdiff_order=5)	
hdiff_w_efdt_ratio	R	15.0		ratio of e-folding time to time step for diffusion	iequations=3
				on vertical wind speed	
hdiff_min_efdt_ratio	$_{ m R}$	1.0		minimum value of hdiff_efdt_ratio near model	iequations=3 .AND.
				top	hdiff_order=4
hdiff_tv_ratio	m R	1.0		Ratio of diffusion coefficients for temperature	
				and normal wind: $T:v_n$	
hdiff_multfac	$_{ m R}$	1.0		Multiplication factor of normalized diffusion	n_dom>1
				coefficient for nested domains	
hdiff_smag_fac	$_{ m R}$	0.15		Scaling factor for Smagorinsky diffusion	iequations=3
		(hydro)			
		0.015			
		(NH)			

Defined and used in: src/namelists/mo_diffusion_nml.f90

4.4.3 dynamics_nml

This namelist is relevant if run_nml:ldynamics=.TRUE.

<u></u>	E	,			7	Γ
Parameter	Type	Default Unit	Unit	Description	Scope	
iequations	I	3		Equations and prognostic variables. Use		
				positive indices for the atmosphere and		
				negative indices for the ocean.		
				0: shallow water model		
	-			1: hydrostatic atmosphere, T		
				2: hydrostatic atm., θ -dp		
				3: non-hydrostatic atmosphere		
				-1: hydrostatic ocean		

Parameter	Type	Default Unit	Unit	Description S	Scope
div_method	I	1		Method for divergence computation:	
				1: Standard Gaussian integral.	
				Hydrostatic atm. model: for unaveraged normal	
				components	
				Non-hydrostatic atm. model: for averaged	
				normal components	
				2: bilinear averaging of divergence	
divavg-cntrwgt	R	0.5		Weight of central cell for divergence averaging id	$idiv_method = 2$
lcoriolis	Γ	TRUE.		Coriolis force	
sw_ref_height	R	*6.0	m	Reference height of shallow water model used	
		$2.94\mathrm{e}4/\mathrm{g}$		for linearization in the semi-implicit time	
				stepping scheme	

Defined and used in: src/namelists/mo_dynamics_nml.f90

4.4.4 echam_conv_nml

Parameter	Type	Default	Unit	Description	Scope
iconv	I	1		Choice of cumulus convection scheme.	iforcing $= 2$.AND.
				1: Nordeng scheme	lconv = .TRUE.
				2: Tiedtke scheme	
				3: hybrid scheme	
ncvmicro	I	0		Choice of convective microphysics scheme.	iforcing $= 2$.AND.
					lconv = .TRUE.
lmfpen	Г	TRUE.		Switch on penetrative convection.	iforcing $= 2$.AND.
					lconv = .TRUE.
lmfmid	Г	TRUE.		Switch on midlevel convection.	iforcing $= 2$.AND.
					lconv = .TRUE.
lmfdd	Г	TRUE.		Switch on cumulus downdraft.	iforcing = 2 .AND.
					lconv = .TRUE.
lmfdudv	Г	TRUE.		Switch on cumulus friction.	iforcing $= 2$.AND.
					lconv = .TRUE.

Parameter	Type	Default Unit	Unit	Description	Scope
cmftau	${f R}$	10800.		Characteristic convective adjustment time	iforcing $= 2$.AND.
				scale.	lconv = .TRUE.
cmfctop	R	0.3		Fractional convective mass flux (valid range	iforcing $= 2$.AND.
				[0,1]) across the top of cloud	lconv = .TRUE.
cprcon	\mathbf{R}	1.0e-4		Coefficient for determining conversion from	iforcing $= 2$.AND.
				cloud water to rain.	lconv = .TRUE.
cminbuoy	\mathbf{R}	0.025		Minimum excess buoyancy.	iforcing $= 2$.AND.
					lconv = .TRUE.
entrpen	\mathbf{R}	1.0e-4		Entrainment rate for penetrative convection.	iforcing $= 2$.AND.
					lconv = .TRUE.
dlev	R	3.e4	Pa	Critical thickness necessary for the onset of	iforcing $= 2$.AND.
				convective precipitation.	lconv = .TRUE.

Defined and used in: src/namelists/mo_echam_conv_nml.f90

4.4.5 ensemble_pert_nml

Parameter	Type	Default	Unit	Description	Scope	
use_ensemble_pert	Г	FALSE.		Main switch to activate physics parameter	run_nml:iforcing =	
				perturbations for ensemble forecasts /	inwp	
				ensemble data assimilation; the perturbations		
				are applied via random numbers depending on		
				the perturbationNumber (ensemble member		
				ID) specified in gribout_nml		
range-gkwake	R	0.333		Variability range for low level wake drag		
				constant		
range-gkdrag	R	0.04		Variability range for orographic gravity wave		
				drag constant		
range-gfluxlaun	\mathbf{R}	0.75e-3		Variability range for non-orographic gravity		
				wave launch momentum flux		
range_zvz0i	R	0.2	s/m	Variability range for terminal fall velocity of ice \mid inwp-gscp = 1 or 2	inwp_gscp = 1 or 2	

Parameter	Type	Default	Unit	Description	Scope
range_entrorg	R	0.2e-3	1/m	Variability range for entrainment parameter in convection scheme	inwp_convection = 1
range_capdcfac_et	R	0.75		Maximum fraction of CAPE diurnal cycle correction applied in the extratropics	icapdcycl = 3
range_rhebc	R	0.05		Variability range for RH threshold for the onset of evaporation below cloud base	inwp_convection = 1
range_texc	뀖	0.05	X	Variability range for temperature excess value in test parcel ascent	inwp_convection = 1
range-box-liq	씸	0.01		Variability range for box width scale of liquid clouds in cloud cover scheme	$invp_cldcover = 1$
range_tkhmin	R	0.2		Variability range for minimum vertical diffusion for heat/moisture	$invp_turb = 1$
range_tkmmin	R	0.2		Variability range for minimum vertical diffusion for momentum	$inwp_turb = 1$
range_tkred_sfc	R	4.0		Range for multiplicative change of reduction of minimum diffusion coefficients near the surface	$inwp_turb = 1$
range_rlam_heat	R	3.0		Variability range (multiplicative!) of laminar transport resistance parameter	$inwp_turb = 1$
range_charnock	R	1.5		Variability range (multiplicative!) of upper and lower bound of wind-speed dependent Charnock parameter	$inwp_turb = 1$
range_minsnowfrac	R	0.05		Variability range for minimum value to which snow cover fraction is artificially reduced in case of melting snow	idiag_snowfrac = $20/30/40$
range_z0_lcc	R	0.25		Variability range (relative change) of roughness length attributed to each landuse class	
range_rootdp	R	0.2		Variability range (relative change) of root depth attributed to each landuse class	
range_rsmin	~	0.2		Variability range (relative change) of minimum stomata resistance attributed to each landuse class	

Parameter	Type	Default	Unit	Description	Scope
range_laimax	R	0.15		Variability range (relative change) of leaf area	
				index (maximum of annual cycle) attributed to	
				each landuse class	

Defined and used in: src/namelists/mo_ensemble_pert_nml.f90

4.4.6 gribout_nml

Scope	filetype=2							filetype=2	filetype=2		filetype=2					filetype=2					
Description	Setting this different to "none" enables a	couple of defaults for the other gribout_nml	namelist parameters. If, additionally, the user	tries to set any of these other parameters to a	conflicting value, an error message is thrown.	Possible values are "none", "deterministic",	"ensemble".	Main switch for Table version	Background process	- GRIB2 code table backgroundProcess.table	Output generating center. If this key is not set,	center information is taken from the grid file	DWD: 78	MPIMET: 98	ECMWF: 98	Output generating Subcenter. If this key is not	set, subcenter information is taken from the	grid file	DWD: 255	MPIMET: 232	ECMWF: 0
Unit	".																				
Default	"determ							15	0		-1					-1					
Type	C							Ι	I		I					I					
Parameter	preset							tablesVersion	backgroundProcess		generatingCenter					generatingSubcenter					

Scope	tifier fletype=2		ifier.table	Local definiton for ensemble products, (only set filetype=2	efault)	mble products, (only set \mid filetype=2	efault)	ta filetype=2		time fletyne=2			mble products (only set filetype=2	efault)	cess filetype=2		filetype=2		r filetype=2			ment filetype=2		mble products (only set \mid filetype=2	efault)	or invariant and $ $ filetype $= 2$		-01-01,	-01-01,	-01-01,	-01-01,
generating Process Identifier		- GRIB2 code table	generatingProcessIdentifier.table	Local definiton for ensem	if value changed from default)	Local definition for ensemble products, (only set	if value changed from default)	Production status of data	- GRIB2 code table 1.3	Significance of reference time	- GRIB2 code table 1.2		Local definition for ensemble products (only set	if value changed from default)	Type of generating process	- GRIB2 code table 4.3	Type of data	- GRIB2 code table 1.4	local Definition Number	- GRIB2 code table	grib2LocalSectionNumber.78.table	local Number of Experiment		Local definition for ensemble products (only set	if value changed from default)	Special reference date for invariant and	climatological fields	TRUE.: set special reference date 0001-01-01,	TRUE.: set special refer	TRUE.: set special refer 00:00	TRUE.: set special refer 00:00
Onit																															
Default	1			-1		-1		1			ı		-1		-1		<u> </u>		<u> </u>			1		-		FALSE.					
$_{\mathrm{Type}}$	I(n-dom)			Ι		I		I		_			I		Ι		Ι		Ι			ш		щ		Γ					
Parameter	generatingProcess	Identifier		numberOfForecastsIn-	Ensemble	perturbationNumber		productionStatusOfPro-		cesseaData significanceOfReference-	0	Time	type Of Ensemble Forecast		typeOfGeneratingPro-	cess	typeOfProcessedData		localDefinitionNumber			localNumberOfExperi-	ment	localTypeOfEnsemble-	Forecast	lspecialdate_invar					

Parameter	Type	Default	Unit	Default Unit Description	Scope
ldate_grib_act	Г	TRUE.		GRIB creation date	filetype=2
				.TRUE.: add creation date	
				.FALSE.: add dummy date	
lgribout_24bit	Γ	FALSE.		If TRUE, write thermodynamic fields ρ , θ_v , T ,	filetype=2
				p with 24bit precision instead of 16bit	

Defined and used in: src/namelists/mo_gribout_nml.f90

4.4.7 grid_nml

Parameter	Type	Default	Unit	Description	Scope
cell_type	I	3		Cell type: not used	
Iplane	L	FALSE.		planar option	
is_plane_torus	L	FALSE.		f-plane approximation on triangular grid	
corio_lat	R	0.0	deg	Center of the f-plane is located at this	lplane=.TRUE. and
				geographical latitude	is_plane_torus=.TRUE.
grid_angular_velocity	R	Earth's	rad/s	The angular velocity in rad per sec.	
l_limited_area	Γ	FALSE.			
grid_rescale_factor	R	1.0		Defined as the inverse of the reduced-size earth	
				reduction factor X . Choose	
				grid_rescale_factor < 1 for a reduced-size	
				earth.	
				The geometry and the timestep will be	
				multiplied by this factor.	
				The angular velocity will be divided by this	
				factor.	
lfeedback	$ L(n_{-}dom) $.TRUE.	.TRUE.		Specifies if feedback to parent grid is	n_dom>1
				performed. Setting lfeedback (1) =.false. turns	
				off feedback for all nested domains; to turn off	
				feedback for selected nested domains, set	
				leedback(1)=.true. and set ".false." for the	
				desired model domains	

Parameter	Type	Default	Unit	Description	Scope
ifeedback_type	I	2		1: incremental feedback	n_dom>1
				2: relaxation-based feedback	
				Note: vertical nesting requires option 2 to run	
				numerically stable over longer time periods	
start_time	$R(n_{-dom})$	0.	w	Time when a nested domain starts to be active	n_dom>1
				(namelist entry is ignored for the global	
				domain)	
end_time	$R(n_{-}dom)$	1.E30	w	Time when a nested domain terminates	n_dom>1
				(namelist entry is ignored for the global	
				domain)	
patch_weight	$ $ R(n_dom)	0.		If patch_weight is set to a value > 0 for any of	n_dom>1
				the first level child patches, processor splitting	
				will be performed, i.e. every of the first level	
				child patches gets a subset of the total number	
				or processors corresponding to its	
				patch_weight. A value of 0. corresponds to	
				exactly 1 processor for this patch, regardless of	
				the total number of processors. For the root	
				patch and higher level childs, patch_weight is	
				not used. However, patch_weight must be set to	
				0 for these patches to avoid confusion.	
lredgrid_phys	ı	.FALSE.		If set to .true. radiation is calculated on a	
				reduced grid (= one grid level higher)	
${\bf dynamics_grid_}$	C			Array of the grid filenames to be used by the	
filename				dycore. May contain the keyword <path></path>	
				which will be substituted by model_base_dir.	
dynamics_parent_	I(n-dom)	i-1		Array of the indexes of the parent grid	
grid_id				filenames, as described by the	
				dynamics-grid-filename array. Indexes start at	
				1, an index of 0 indicates no parent.	

Parameter	Type	Default	Unit	Description Sc	Scope
radiation_grid_	C			Array of the grid filenames to be used for the lre	lredgrid_phys=.TRUE.
filename				radiation model. Filled only if the radiation	
				grid is different from the dycore grid. May	
				contain the keyword <pre><pre><pre>contain the keyword <pre><pre><pre>path></pre> which will be</pre></pre></pre></pre></pre>	
				substituted by model_base_dir.	
dynamics_radiation_g	I(n-dom)	1 for i=1		Array of the indexes linking the dycore grids,	
rid_link				as described by the dynamics-grid_filename	
				array, and the radiation-grid-filename array. It	
				provides the link index of the	
				radiation_grid_filename, for each entry of the	
				dynamics-grid_filename array. Indexes start at	
				1, an index of 0 indicates that the radiation	
				grid is the same as the dycore grid. Only needs	
				to be filled when the radiation_grid_filename is	
				defined.	
create_vgrid	Γ	.FALSE.		.TRUE.: Write vertical grid files containing	
				(vct_a, vct_b, z_ifc, and z_ifv.	
vertical_grid_filename	$C(n_{-dom})$			Array of filenames. These files contain the	
				vertical grid definition (vct_a, vct_b, z_ifc). If	
				empty, the vertical grid is created within ICON	
				during the setup phase.	
use_duplicated_	Γ	TRUE.		if .TRUE., the zero connectivity is replaced by	
connectivity				the last non-zero value	
use_dummy_cell_closure	J	FALSE.		if .TRUE. then create a dummy cell and	
				connect it to cells and edges with no neighbor	

Defined and used in: src/namelists/mo_grid_nml.f90

4.4.8 gridref_nml

Parameter	Type	Default	Unit	Description	Scope
grf_intmethod_c	I	2		Interpolation method for grid refinement	n_dom>1
				(cell-based dynamical variables):	
				2: gradient-based interpolation	
grf_intmethod_ct	I	2		Interpolation method for grid refinement	n_dom>1
				(cell-based tracer variables):	
				1: parent-to-child copying	
				2: gradient-based interpolation	
$\operatorname{grf_intmethod_e}$	I	9		Interpolation method for grid refinement	n_dom>1
				(edge-based variables):	
				1: inverse-distance weighting (IDW)	
				2: RBF interpolation	
				3: combination gradient-based / IDW	
				4: combination gradient-based / RBF	
				5/6: same as $3/4$, respectively, but direct	
				interpolation of mass fluxes along nest interface	
				edges	
grf_velfbk	I	1		Method of velocity feedback:	n_dom>1
				1: average of child edges 1 and 2	
				2: 2nd-order method using RBF interpolation	
grf_scalfbk	I	2		Feedback method for dynamical scalar	n_dom>1
				variables (T, p_{sfc}) :	
				1: area-weighted averaging	
				2: bilinear interpolation	
grf_tracfbk	Ι	2		Feedback method for tracer variables:	n_dom>1
				1: area-weighted averaging	
				2: bilinear interpolation	
grf_idw_exp_e12	R	1.2		exponent of generalized IDW function for child	n_dom>1
				edges 1/2	
grf_idw_exp_e34	R	1.7		exponent of generalized IDW function for child	n-dom>1
				edges $3/4$	
rbf_vec_kern_grf_e	_			RBF kernel for grid refinement (edges):	n_dom>1
		_		I: Gaussian	

Scope		n_dom>1						n_dom>1		$n_dom > 1$		n_dom>1		$n_{-}dom > 1$.AND.	lfeedback = .TRUE.	n_dom;1 .AND.	leedback = .TRUE.	.AND. ifeedback_type	= 2
Description	$2: 1/(1+r^2)$ 3: inverse multiquadric	RBF scale factor for grid refinement (lateral	boundary interpolation to edges). Refers to the	respective parent domain and thus does not	need to be specified for the innermost nest.	Lower values than the default of 0.5 are needed	for child mesh sizes less than about 500 m.	Deniminator for lateral boundary diffusion of	temperature	Deniminator for lateral boundary diffusion of	velocity	.TRUE.: Apply mass conservation correction in	feedback routine	TRUE:: Apply density nudging near lateral	nest boundary if grf_intmethod_e ≤ 4	Relaxation time scale for feedback			
Unit																			
Default		0.5						135		200		FALSE.		FALSE.		10800			
Type		$R(n_{-dom})$						R		\mathbf{R}		Γ		Γ		R			
Parameter		$ m rbf_scale_grf_e$						denom_diffu_t		denom_diffu_v		l_mass_consvcorr		l_density_nudging		fbk_relax_timescale			

Defined and used in: src/namelists/mo_gridref_nml.f90

4.4.9 gw_hines_nml (Scope: Igw_hines = .TRUE. in echam_phy_nml)

	[7
Parameter	Type	Default	Unit	Unit Description	Scope
Iheatcal	Г	.FALSE.		.TRUE.: compute drag, heating rate and	
				diffusion coefficient from the dissipation of	
				gravity waves	
				.FALSE.: compute drag only	

Scope																	$lrmscon_lat = .TRUE.$	$lrmscon_lat = .TRUE.$	$lrmscon_lat = .TRUE.$
Description	Index of model level, counted from the surface,	from which the gravity wave spectra are	emitted	Root mean square gravity wave wind at the	emission level	Typical gravity wave horizontal wavenumber	Minimum bound in vertical wavenumber	.TRUE.: use latitude dependent rms wind	-—latitude— >= lat_rmscon: use rmscon	- —latitude— <= lat_rmscon_eq: use	rmscon_eq	- lat_rmscon_eq < —latitude— < lat_rmscon:	use linear interpolation between rmscon_eq and	rmscon	.FALSE.: use globally constant rms wind	rmscon	rmscon_eq is used equatorward of this latitude	rmscon is used polward of this latitude	is used equatorward of latitude lat_rmscon_eq
Unit				s/m		1/m	1/m										deg N	deg N	m/s
Default	10			1.0		5.0e-5	0.0	FALSE.									5.0	10.0	1.2
Type	П			$_{\rm R}$		R	R	Г									$_{\rm R}$	R	\mathbf{R}
Parameter	emiss_lev			rmscon		kstar	m-min	lrmscon_lat									lat_rmscon_eq	lat_rmscon	rmscon_eq

Defined and used in: src/namelists/mo_gw_hines_nml.f90

4.4.10 ha_dyn_nml

This namelist is relevant if run_nml:ldynamics=.TRUE. and dynamics_nml:iequations=IHS_ATM_TEMP or IHS_ATM_THETA.

Parameter	$\mid \mathrm{Type} \mid$	Default	Unit	Pefault Unit Description	Scope
itime_scheme	I	14		Time integration scheme:	
				11: pure advection (no dynamics)	
				12: 2 time level semi implicit (not yet	
				implemented)	
				13: 3 time level explicit	

ileapfrog_startup I	1 0.1		14: 3 time level with semi implicit correction	
ileapfrog-startup	1 0.1		15. Atomorphy Ath and man Dimer It with another	_
ileapfrog_startup I	1 0.1		19: Standard 4th-order runge-rutta method	
ileapfrog_startup I	1 0.1		(4-stage)	
ileapfrog_startup I	1 0.1		16: SSPRK(5,4) scheme $(5-stage)$	
	0.1		How to integrate the first time step when the	itime_scheme= $13 \text{ or } 14$
	0.1		leapfrog scheme is chosen. $1 = \text{Euler forward}$; 2	
	0.1		= a series of sub-steps.	
asselin_coeff R			Asselin filter coefficient	itime_scheme= $13 \text{ or } 14$
si_2tls R	9.0		weight of time step $n+1$. Valid range: $[0,1]$	itime_scheme=12
si_expl_scheme I	2		scheme for the explicit part used in the 2 time	$itime_scheme=12$
			level semi-implicit time stepping scheme. $1 =$	
			Euler forward; $2 = Adams$ -Bashforth 2nd order	
si_cmin R	30.0	m/s	semi implicit correction is done for eigenmodes	itime_scheme=14 and
			with speeds larger than si_cmin	lsi_3d=.FALSE.
	1.0		weight of the semi implicit correction	itime_scheme=14
si_offctr R	0.7			itime_scheme=14
R	1.0e-3		relative tolerance for GMRES solver	itime_scheme=14
J	FALSE.		3D GMRES solver or decomposistion into 2D	lshallow_water=.FALSE.
			problems	and itime_scheme=14
ldry_dycore L	TRUE.		Assume dry atmosphere	$iequations \in \{1,2\}$
lref_temp L	.FALSE.		Set a background temperature profile as base	$iequations \in \{1,2\}$
			state when computing the pressure gradient	
			force	

4.4.11 initicon_nml

Parameter	Type	Default	Unit	Description	Scope
init_mode init_mode	E H	10800		1: MODE_DWDANA start from DWD analysis or FG 2: MODE_IFSANA start from IFS analysis 3: MODE_COMBINED IFS atm + ICON/GME soil 4: MODE_COSMODE start from COSMO-DE forecast 5: MODE_IAU analysis update. Extension of MODE_IAU_OLD start from DWD analysis with increments 6: MODE_IAU_OLD start from DWD analysis with increments 6: MODE_IAU_OLD start from DWD analysis with incremental analysis update. NOTE: Extension of mode MODE_DWDANA_INC including W_SO increments. 7: MODE_ICONVREMAP start from DWD first guess with subsequent vertical remapping (work in progress; so far, changing the number of model levels does not yet work) Time interval during which an incremental	init_mode=5,6
dt_shift	R	0	∞	analysis update (IAU) is performed Time by which the actual model start time is	init_mode=5,6
iterate_iau	Ţ	.FALSE.		shifted ahead of the nominal date. Must be NEGATIVE, usually -0.5 dt_iau. If .TRUE., the IAU phase is calculated twice with halved dt_shift in first cycle (allows writing a fully initialized analysis at the nominal initialization date while using a centered IAU window for the forecast).	init_mode=5,6 and dt_shift < 0

Parameter	Type	Default	Unit	Description	Scope
start_time_avg_fg	R	0	w	Start time for calculating temporally averaged	4
				first guess output for data assimilation.	
end_time_avg_fg	R	0	x	End time for calculating temporally averaged	
				first guess output for data assimilation.	
				Setting end_time_avg_fg > start_time_avg_fg	
				activates the averaging	
interval_avg_fg	R	0	œ	Corresponding averaging interval. Note that	
				end_time_avg_fg — start_time_avg_fg must not	
				be smaller than the averaging interval	
rho_incr_filter_wgt	R	0		Vertical filtering weight on density increments	init_mode=5,6
niter_diffu	Ι	10		Number of diffusion iterations applied on wind	init_mode=5,6
				increments	
niter_divdamp	I	25		Number of divergence damping iterations	init_mode=5,6
				applied on wind increments	
type_iau_wgt	Ι	П		Weighting function for performing IAU	init_mode=5,6
				1: Top-Hat	
				2: SIN2	
nlevsoil_in	I	4		number of soil levels of input data	init_mode=2
zpbl1	\mathbf{R}	500.0	m	bottom height (AGL) of layer used for gradient	
				computation	
zpbl2	$_{\rm R}$	1000.0	m	top height (AGL) of layer used for gradient	
	,	[computation	
lread_ana	7	TRUE.		If .FALSE., ICON is started from first guess	$init_mode=1,3$
				only. Analysis field is not required, and skipped	
				if provided.	
lconsistency_checks	Г	TRUE.		If .FALSE., consistency checks for Analysis and	$init_mode=1,3,4,5,6$
				First Guess fields are skipped. On default,	
				checks are performed for $uuidOfHGrid$ and	
				validity time.	
l_coarse2fine_mode	L(n-dom)	FALSE.		If true, apply corrections for coarse-to-fine	
				mesn interpolation to wind and temperature	

Parameter	Type	Default	Unit	Description	Scope
lp2cintp_incr	L(n-dom)	.FALSE.		If true, interpolate atmospheric data	init_mode=5,6
	,			assimilation increments from parent domain.	
				Can be specified separately for each nested	
				domain; setting the first (global) entry to true	
				activates the interpolation for all nested	
				domains.	
lp2cintp_sfcana	L(n-dom)	FALSE.		If true, interpolate atmospheric surface analysis	init_mode=5,6
				data from parent domain.	
				Can be specified separately for each nested	
				domain; setting the first (global) entry to true	
				activates the interpolation for all nested	
				domains.	
ltile_init	L	FALSE.		True: initialize tiled surface fields from a first	$init_mode=1,5,6$
				guess coming from a run without tiles.	
				Along coastlines and lake shores, a neighbor	
				search is executed to fill the variables on	
				previously non-existing land or water points	
				with reasonable values. Should be combined	
				with ltile_coldstart = .TRUE.	
ltile_coldstart	J	FALSE.		If true, tiled surface fields are initialized with	$init_mode=1,5,6$
				tile-averaged fields from a previous run with	
				tiles.	
				A neighbor search is applied to subgrid-scale	
				ocean points for SST and sea-ice fraction.	
lvert_remap_fg	ı	FALSE.		If true, vertical remapping is applied to the	init_mode=5,6
				atmospheric first-guess fields, whereas the	
				analysis increments remain unchanged. The	
				number of model levels must be the same for	
				input and output fields, and the z_ifc (alias	
				HHL) field pertaining to the input fields must	
				be appended to the first-guess file.	

Parameter	Type	Default	Unit	Description	Scope
ifs2icon_filename	C			Filename of IFS2ICON input file, default	init_mode=2
				" <path>ifs2icon_R<nroot>B<jlev>_DOM</jlev></nroot></path>	
				<pre><idom>.nc". May contain the keywords <path></path></idom></pre>	
				which will be substituted by model_base_dir,	
				as well as nroot, jlev, and idom defining the	
				current patch.	
dwdfg_filename	C			Filename of DWD first-guess input file, default	$init_mode=1,3,5,6$
				" <path>dwdFG_R<nroot>B<jlev>_DOM</jlev></nroot></path>	
				<idom>.nc". May contain the keywords <pre><pre>class</pre></pre></idom>	
				which will be substituted by model_base_dir,	
				as well as nroot, jlev, and idom defining the	
				current patch.	
dwdana_filename	C			Filename of DWD analysis input file, default	$init_mode=1,3,5,6$
				" <path>dwdana_R<nroot>B<jlev>_DOM</jlev></nroot></path>	
				<idom>.nc". May contain the keywords <path></path></idom>	
				which will be substituted by model_base_dir,	
				as well as nroot, jlev, and idom defining the	
				current patch.	
filetype	I	-1		One of CDI's FILETYPE_XXX constants.	
		(undef.)		Possible values: 2 (=FILETYPE_GRB2), 4	
				(=FILETYPE_NC2). If this parameter has not	
				been set, we try to determine the file type by	
				its extension "*.grb*" or ".nc".	
ana_varlist	C(:)			List of mandatory analysis fields for the global	$init_mode=1,5,6$
				domain that must be present in the analysis	
				file. If these fields are not found, the model	
				aborts. For all other analysis fields, the	
				FG-fields will serve as fallback position.	
ana_varlist_n2	C(:)			List of mandatory analysis fields for domain 2	init_mode=5,6
				that must be present in the analysis file. If	
				these fields are not found, the model aborts.	
				For all other analysis fields, the FG-fields will	
				serve as fallback position.	

Parameter	Type	Default	Unit	Default Unit Description	Scope
ana_varnames_map_	C			Dictionary file which maps internal variable	
file				names onto GRIB2 shortnames or NetCDF var	
				names. This is a text file with two columns	
				separated by whitespace, where left column:	
				ICON variable name, right column: GRIB2	
				short name.	

Defined and used in: src/namelists/mo_initicon_nml.f90

4.4.12 interpol_nml

Parameter	Type	Default	Unit	Description	Scope
l_intp_c2l	Г	.TRUE.		DEPRECATED	
l_mono_c2l	l l	TRUE.		Monotonicity can be enforced by demanding	
				that the interpolated value is not higher or	
				lower than the stencil point values.	
llsq_high_consv	ı	TRUE.		conservative (T) or non-conservative (F)	
				least-squares reconstruction for high order	
				transport	
lsq_high_ord	П	3		polynomial order for high order reconstruction	
				1: linear	ihadv_tracer=4
				2: quadratic	
				30: cubic (no 3^{rd} order cross deriv.)	
				3: cubic	
llsq_lin_consv	ı	.FALSE.		conservative (T) or non-conservative (F)	
				least-squares reconstruction for 2nd order	
				(linear) transport	
nudge_efold_width	$_{ m R}$	2.0		e-folding width (in units of cell rows) for lateral	
				boundary nudging coefficient	
nudge_max_coeff	${ m R}$	0.02		Maximum relaxation coefficient for lateral	
				boundary nudging	

Parameter	Type	Default	Unit	Description	Scope
nudge_zone_width	I	∞		Total width (in units of cell rows) for lateral	
				boundary nudging zone. If i 0 the patch boundary-depth index is used.	
${ m rbf_dim_c2l}$	Ι	10		stencil size for direct lon-lat interpolation: $4 =$	
				nearest neighbor, $13 = \text{vertex stencil}$, $10 =$	
				edge stencil.	
rbf_scale_mode_ll	П	2		Specifies, how the RBF shape parameter is	
				determined for lon-lat interpolation.	
				1: lookup table based on grid level	
				2 : determine automatically.	
				So far, this routine only estimates the smallest	
				value for the shape parameter for which the	
				Cholesky is likely to succeed in floating point	
				arithmetic. 3: explicitly set shape parameter in	
				each output namelist (namelist parameter	
				output_nml::rbf_scale, p. 109).	
rbf_vec_kern_c	I	1		Kernel type for reconstruction at cell centres:	
				1: Gaussian	
				3: inverse multiquadric	
rbf_vec_kern_e	Н	3		Kernel type for reconstruction at edges:	
				1: Gaussian	
				3: inverse multiquadric	
rbf_vec_kern_ll	П			Kernel type for reconstruction at lon-lat-points:	
				1: Gaussian	
				3: inverse multiquadric	
rbf_vec_kern_v	П			Kernel type for reconstruction at vertices:	
				1: Gaussian	
				3: inverse multiquadric	
rbf_vec_scale_c	$R(n_{-dom})$	resolution-		Scale factor for RBF reconstruction at cell	
				centres	
rbf_vec_scale_e	$R(n_{-}dom)$			Scale factor for RBF reconstruction at edges	
		dependen			

Default Unit Description Scope	Scale factor for RBF reconstruction at vertices		Flag. If .FALSE. barycentric interpolation is	replaced by a fallback interpolation.
Unit				
Default	resolution-	dependent	FALSE.	
Type	$R(n_{-dom})$		L	
Parameter	rbf_vec_scale_v		support_baryctr_intp	

Defined and used in: src/namelists/mo_interpol_nml.f90

4.4.13 io_nml

lkeep in_sync L .FALSE. dt_diag R 86400. s dt_checkpoint R 2592000 s inextra_2d I 0 inextra_3d I 0			2
Action R 86400. R 2592000 I 0 I 0		Sync output stream with file on disk after each	
Doint R 86400. I 0 I 0		timestep	
Doint R 2592000 I 0 I 0	w	diagnostic integral output interval	run_nml:output =
Doint R. 2592000			"totint"
inextra_2d I 0 inextra_3d I 0	œ	Time interval for writing restart files. Note that	output /= "none"
inextra_2d I 0 inextra_3d I 0		if the value of dt_checkpoint resulting from	(run_nml)
inextra_2d I 0 inextra_3d I 0		model default or user's specification is longer	
inextra_2d I 0 inextra_3d I 0		than time_nml:dt_restart, it will be reset (by	
inextra_2d I 0 inextra_3d I 0		the model) to dt_restart so that at least one	
inextra_2d I 0 inextra_3d I 0		restart file is generated during the restart cycle.	
inextra_3d I 0		Number of extra 2D Fields for	dynamics_nml:iequations
inextra_3d I 0		diagnostic/debugging output.	= 3 (to be done for 1,
inextra_3d I 0			2)
		Number of extra 3D Fields for	dynamics_nml:iequations
		diagnostic/debugging output.	= 3 (to be done for 1,
			2)
lflux_avg L .TRUE.		if .FALSE. the output fluxes are accumulated	iequations=3
		from the beginning of the run	iforcing=3
		if .TRUE. the output fluxes are average values	
		from the beginning of the run, except of TOT PREC that would be accumulated	

Parameter	Type	Default	Unit	Description	Scope
itype-pres-msl	1	1		Specifies method for computation of mean sea level pressure (and geopotential at pressure levels below the surface). 1: GME-type extrapolation, 2: stepwise analytical integration, 3: current IFS method, 4: IFS method with consistency correction 5: New DWD method constituting a mixture between IFS and old GME method (departure level for downward extrapolation between 10 m and 150 m AGL depending on elevation)	
itype_rh	I	1		Specifies method for computation of relative humidity 1: WMO-type: water only (e_s=e_s_water), 2: IFS-type: mixed phase (water and ice), 3: IFS-type with clipping (rh \le 100)	
gust_interval output_nml_dict	면 D	3600.	ω	Interval over which wind gusts are maximized File containing the mapping of variable names to the internal ICON names. May contain the keyword <pre>cpath> which will be substituted by model_base_dir.</pre> The format of this file: One mapping per line, first the name as given in the ml_varlist, hl_varlist, pl_varlist or il_varlist of the output_nml namelists, then the internal ICON name, separated by an arbitrary number of blanks. The line may also start and end with an arbitrary number of blanks. Empty lines or lines starting with # are treated as comments. Names not covered by the mapping are used as they are.	iforcing=3 output_nml namelists

Parameter	Type	Default	Unit	Description	Scope
netcdf_dict	C	, ,		File containing the mapping from internal	output_nml namelists,
				names to names written to NetCDF. May	NetCDF output
				contain the keyword <pre><pre>contain the keyword <pre><pre>contain</pre></pre></pre></pre>	
				substituted by model_base_dir.	
				The format of this file:	
				One mapping per line, first the name written	
				to NetCDF, then the internal name, separated	
				by an arbitrary number of blanks (inverse to	
				the definition of output_nml_dict). The line	
				may also start and end with an arbitrary	
				number of blanks. Empty lines or lines starting	
				with $\#$ are treated as comments.	
				Names not covered by the mapping are output	
				as they are.	
				Note that the specification of output variables,	
				e.g. in ml_varlist, is independent from this	
				renaming, see the namelist parameter	
				output_nml_dict for this.	
lnetcdf_flt64_output	Γ	.FALSE.		If .TRUE. floating point variable output in	
				NetCDF files is written in 64-bit instead of	
				32-bit accuracy.	
				This is currently implemented for the atm.	
				dynamical core and ECHAM physics.	
restart_file_type	I	4		Type of restart file. One of CDI's	
				FILETYPE_XXX. So far, only 4	
				(=FILETYPE_NC2) is allowed	
lmask_boundary	Γ	ĹΉ		Set to .TRUE., if interpolation zone should be	
				masked in output.	

Defined and used in: src/namelists/mo_io_nml.f90

4.4.14 les_nml (parameters for LES turbulence scheme; valid for inwp_turb=5)

Parameter	Type	Default	Unit	Description	Scope
sst	R	300	K	sea surface temperature for idealized LES	isrfc_type=5,4
				simulations	
shfix	$_{ m R}$	0.1	$\mathrm{Km/s}$	Kinematic sensible heat flux at surface	$isrfc_{-type} = 2$
lhflx	R	0	m/s	Kinematic latent heat flux at surface	$isrfc_{type} = 2$
srfc_type	I	1		surface type	
				0 = No fluxes and zero shear stress	
				1 = TERRA land physics	
				2 = fixed surface fluxes	
				3 = fixed buoyancy fluxes	
				4 = RICO test case	
				5 = fixed SST	
ufric	R	666-	s/m	friction velocity for idealized LES simulations;	
				if i 0 then it is automatically diagnosed	
psfc	R	-666	Pa	surface pressure for idealized LES simulations;	
				if i 0 then it uses the surface pressure from	
				dynamics	
min_sfc_wind	R	1.0	s/m	Minimum surface wind for surface layer useful	
				in the limit of free convection	
is_dry_cbl	L	FALSE.		switch for dry convective boundary layer	
				simulations	
smag_constant	$_{ m R}$	0.23		Smagorinsky constant	
km_min	R	0.0		Minimum turbulent viscosity	
max_turb_scale	$_{ m R}$	300.0		Asymtotic maximum turblence length scale	
				(useful for coarse grid LES and when grid is	
				vertically stretched)	
turb_prandtl	Я	0.333333		turbulent Prandtl number	
pflux	R	0.0007	$\mathrm{m}^2/\mathrm{s}^3$	buoyancy flux for idealized LES simulations	$isrfc_type=3$
				(Stevens 2007)	
tran_coeff	R	0.02	m/s	transfer coefficient near surface for idealized	$isrfc_type=3$
				LES simulation (Stevens 2007)	

Parameter	Type	Default Unit	Unit	Description Sc	Scope
vert_scheme_type	I	2		type of time integration scheme in vertical diffusion	
				1 = explicit $2 = fully implicit$	
samp] fred sec	Ω.	09	v.	sampling frequency in seconds for statistical	
Do and Tanana	,)	1	(1D and 0D) output	
avg_interval_sec	R	006	w	(time) averaging interval in seconds for 1D	
				statistical output	
expname	C	ICOLES		expname to name the statistical output file	
ldiag_les_out	L	FALSE.		Control for the statistical output in LES mode	
les_metric	L	.FALSE.		Switch to turn on Smagorinsky diffusion with	
				3D metric terms to account for topography	

Defined and used in: src/namelists/mo_les_nml.f90

4.4.15 limarea_nml (Scope: I_limited_area=.TRUE. in grid_nml)

D		D.f14	TITEST	December	C
rarmeter	Type	Derauit		Description	Scope
itype_latbc	I	0		Type of lateral boundary nudging.	
				0: constant lateral boundary conditions derived	
				from the initial conditions,	
				1: time-dependent lateral boundary conditions	
				provided by an external source (IFS,	
				COSMO-DE or a coarser-resolution ICON	
				run),	
				2: Test mode using time-dependent lateral	
				boundary conditions from a nested ICON run	
				in which the present limited-area domain was	
				operated as a nested grid with identical(!)	
				model level configuration.	

Parameter	Type	Default	Unit	Description	Scope
dtime_latbc	R	10800.0	w	Time difference between two consecutive	itype_latbc ≥ 1
				boundary data.	
init_latbc_from_fg	L	.FALSE.		If .TRUE., take lateral boundary conditions for	itype_latbc ≥ 1
				initial time from first guess (or analysis) field	
nlev_latbc	I	0	w ·	Number of vertical levels in boundary data.	itype_latbc ≥ 1
latbc_filename	C			Filename of boundary data input file, these	itype_latbc ≥ 1
				files must be located in the latbc_path	
				directory. Default:	
				"prepiconR <nroot>B<jlev>_<y><m><d><h>nconR<nroot>nc".</nroot></h></d></m></y></jlev></nroot>	
				The keyword tokens <y>, <m>, <d>, and <h></h></d></m></y>	
				will be automatically replaced during the	
				run-time (year, month, day, hour). In case the	
				time span between two consecutive boundary	
				data is less than 1 hour, one can use <min> and</min>	
				<sec>. The keyword <ddhhmmss> is replaced by</ddhhmmss></sec>	
				a relative day-hour-minute-second string.	
latbc_path	C			Absolute path to boundary data.	itype_latbc ≥ 1
latbc_boundary_grid	C	" "		Grid file defining the lateral boundary. Empty	itype_latbc ≥ 1
				string means: whole domain is read for the	
				lateral boundary. This NetCDF grid file must	
				contain two integer index arrays: int	
				<pre>global_cell_index(cell), int</pre>	
				global_edge_index(edge), both with	
				attributes nglobal which contains the global	
				size size of the non-sparse cells and edges.	

Parameter	Type	Default Unit	Unit	Description S	Scope
latbc_varnames_map_	C			Dictionary file which maps internal variable n	num_prefetch_proc=1
file				names onto GRIB2 shortnames or NetCDF var	
				names. This is a text file with two columns	
				separated by whitespace, where left column:	
				ICON variable name, right column: GRIB2	
				short name. This list contains variables that	
				are to be read asynchronously for boundary	
				data nudging in a HDCP2 simulation. All new	
				boundary variables that in the future, would be	
				read asynchronously. Need to be added to text	
				file dict.latbc in run folder.	

Defined and used in: src/namelists/mo_limarea_nml.f90

4.4.16 Ind_nml

Parameter	Type	Default	Unit	Default Unit Description	Scope
nlev_snow	I	2		number of snow layers	lmulti_snow=.true.
ntiles	I	1		number of tiles	
lsnowtile	Γ	.FALSE.		.TRUE.: consider snow-covered and snow-free	ntiles>1
				tiles separately	
frlnd_thrhld	R	0.05		fraction threshold for creating a land grid point ntiles>1	ntiles>1
frlake_thrhld	R	0.05		fraction threshold for creating a lake grid point	ntiles>1
frsea_thrhld	R	0.05		fraction threshold for creating a sea grid point	ntiles>1
frlndtile_thrhld	R	0.05		fraction threshold for retaining the respective	ntiles>1
				tile for a grid point	
lmelt	Г	TRUE.		.TRUE. soil model with melting process	
lmelt_var	I	TRUE.		.TRUE. freezing temperature dependent on	
				water content	
lana_rho_snow	T	TRUE.		.TRUE. take rho_snow-values from analysis file	init_mode=1
$lmulti_snow$	Γ	FALSE.		TRUE. for use of multi-layer snow model	
				(default is single-sayer scheme)	

Parameter	Type	Default	Unit	Description	Scope
12lay_rho_snow	ı	.FALSE.		.TRUE. predict additional snow density for	$lmulti_snow = .FALSE.$
				upper part of the snowpack, having a	
				maximum depth of max-toplaydepth	
max_toplaydepth	R	0.25	m	maximum depth of uppermost snow layer	lmulti_snow=.TRUE.
					or 12lay rho snow=.TRUE.
idiag_snowfrac	I			Type of snow-fraction diagnosis:	>
)				1 = based on SWE only	
				2-4 = more advanced experimental methods	
				20, 30, 40 = same as 2, 3, 4, respectively, but	
				with artificial reduction of snow fraction in	
				case of melting snow	
itype_lndtbl	I	3		Table values used for associating surface	
				parameters to land-cover classes:	
				1 = defaults from extpar (GLC2000 and	
				GLOBCOVER2009)	
				2 = Tuned version based on IFS values for	
				globcover classes (GLOBCOVER2009 only)	
				3 = even more tuned operational version	
				(GLOBCOVER2009 only)	
				4 = tuned version for new bare soil evaporation	
				scheme (itype_evsl=4)	
itype_root	I	2		root density distribution:	
				1 = constant	
				2 = exponential	
itype_evsl	Ι	2		type of bare soil evaporation parameterization	
				2 = Dickinson (1984)	
				3 = Noilhan and Planton (1989)	
				4 = Resistance-based scheme by Jan-Peter	
				Schulz	

Parameter	Type	Default	Unit	Description S	Scope
itype_heatcond	I	2		type of soil heat conductivity	
				1 = constant soil heat conductivity	
				2 = moisture dependent soil heat conductivity	
				3 = variant of option 2 with reduced	
				near-surface heat conductivity in the presence	
				of plant cover	
itype_interception	I	1		type of plant interception	
				1 = standard scheme, effectively switched off	
				by tiny value cwimax_ml	
				2 = Rain and snow interception (under	
				development)	
cwimax_ml	\mathbf{R}	1.e-6	m	scaling parameter for maximum interception it	itype_interception = 1
				storage (almost switched off);	
				use 5.e-4 to activate interception storage	
c_soil	${ m R}$	1.		surface area density of the (evaporative) soil	
				surface	
				allowed range: $0-2$	
c_soil_urb	\mathbf{R}	1.		surface area density of the (evaporative) soil	
				surface, urban areas	
				allowed range: $0-2$	
itype_hydbound	Ι			type of hydraulic lower boundary condition	
				1 = none	
				3 = ground water as lower boundary of soil	
				column	
Istomata	Γ	TRUE.		If .TRUE., use map of minimum stomatal	
				resistance	
				If .FALSE., use constant value of 150 s/m.	
12tls	Γ	TRUE.		If .TRUE., forecast with 2-Time-Level	
				integration scheme	
lseaice	Γ	TRUE.		TRUE. for use of sea-ice model	
llake	I	TRUE.		.TRUE. for use of lake model	

Parameter	$\mid ext{Type}$	Default	Unit	Description	Scope
sstice_mode	I	1		1: SST and sea ice fraction are read from the	iequations=3
				analysis. The SST is kept constant whereas the	iforcing=3
				sea ice fraction can be modified by the seaice	
				model.	
				2: SST and sea ice fraction are read from the	
				analysis. The SST is updated by climatological	
				increments on a daily basis. The sea ice	
				fraction can be modified by the seaice model.	
				3: SST and sea ice fraction are updated daily,	
				based on climatological monthly means	
				4: SST and sea ice fraction are updated daily,	
				based on actual monthly means	
				5: SST and sea ice fraction are updated daily,	
				based on actual daily means (not yet	
				implemented)	
sst_td_filename	C			Filename of SST input files for time dependent	sstice_mode=2,3
				SST. Default is	
				" <path>SST_<year>_imonth;_<gridfile>".</gridfile></year></path>	
				May contain the keyword <path> which will</path>	
				be substituted by model_base_dir	
ci_td_filename	C			Filename of sea ice fraction input files for time	sstice_mode=2,3
				dependent sea ice fraction. Default is	
				" <path>CI_<year>_<month>_<gridfile>".</gridfile></month></year></path>	
				May contain the keyword <path> which will</path>	
				be substituted by model_base_dir	

Defined and used in: src/namelists/mo_lnd_nwp_nml.f90

4.4.17 Is_forcing_nml (parameters for large-scale forcing; valid for torus geometry)

Parameter	Type	Default	Unit	Default Unit Description	Scope
is_subsidence_moment	Г	FALSE.		switch for enabling LS vertical advection due	is_plane_torus=.TRUE.
				to subsidence for momentum equations	
is_subsidence_heat	Г	FALSE.		switch for enabling LS vertical advection due	is_plane_torus=.TRUE.
				to subsidence for thermal equations	
is_advection	L	FALSE.		switch for enabling LS horizontal advection	is_plane_torus=.TRUE.
				(currently only for thermal equations)	
is_geowind	Γ	FALSE.		switch for enabling geostrophic wind	is_plane_torus=.TRUE.
is_rad_forcing	Γ	FALSE.		switch for enabling radiative forcing	is_plane_torus=.TRUE.
					inwp_rad=.FALSE.
is_theta	Г	FALSE.		switch to indicate that the prescribed radiative	is_plane_torus=.TRUE.
				forcing is for potential temperature	is_rad_forcing=.TRUE.

Defined and used in: src/namelists/mo_ls_forcing_nml.f90

4.4.18 master_model_nml (repeated for each model)

Scope	Character string for naming this component.	the model namelists.		onent to run.					is model.	s model.	
Description	Character string for n	File name containing the model namelists.		Identifies which component to run.	1=atmosphere	2=ocean	3=radiation	99=dummy_model	Start MPI rank for this model.	End MPI rank for this model.	Stride of MPI ranks.
Unit											
Default Unit				7					0	-1	1
Type	C	C		I					П	П	I
Parameter	model_name	model_namelist_	ilename	model_type					nodel_min_rank	nodel_max_rank	model_inc_rank

4.4.19 master_nml

Parameter	Type	Default	Unit	Default Unit Description	Scope
lrestart	Γ	.FALSE.		If .TRUE.: Current experiment is started from	
				a restart.	
lrestart_write_last	Г	.FALSE.		If .TRUE.: model run should create restart at	
				experiment end. This is independent from the	
				settings of the restart interval.	
model_base_dir	C			General path which may be used in file names	
				of other name lists: If a file name contains the	
				keyword " <path>", then this model_base_dir</path>	
				will be substituted.	

4.4.20 meteogram_output_nml

Nearest neighbour 'interpolation' is used for all variables.

Parameter	Type	Default U	Unit	Description Scope	
lmeteogram_enabled	L(n_dom)	.FALSE.		Flag. True, if meteogram of output variables is desired.	
zprefix	C(n_dom)	$C(n_{-}dom)$ "METEO GRAM."		string with file name prefix for output file	
ldistributed	$L(n_{-dom})$	TRUE.		Flag. Separate files for each PE.	
loutput_tiles	ı	FALSE.		Write tile-specific output for some selected surface/soil fields	
$^{ m n0-mtgrm}$	I(n-dom)	0		initial time step for meteogram output.	
ninc_mtgrm	I(n-dom)	1		output interval (in time steps)	
stationlist_tot		53.633,		list of meteogram stations (triples with lat, lon,	
		9.983,		name string)	
		'Ham-			
		burg'			

Parameter	Type	Default Unit I	Unit	Description	Scope
var_list	C(:)	""		Positive-list of variables (optional). Only	
				variables contained in this list are included in	
				the meteogram. If the default list is not	
				changed by user input, then all available	
				variables are added to the meteogram	

Defined and used in: src/namelists/mo_mtgrm_nml.f90

4.4.21 mpi_phy_nml

controlling how the atmospheric boundary conditions for the MPI physics are determined. Time control parameters are available for the The MPI physics is configured by a data structure $mpi_phy_config(jg=1:ndom)\%[param_i]$, which is a 1-dimensional array extending over all domains. The structure contains several parameters providing time control for atmospheric forcings and additional logical switches following atmospheric processes:

cumulus convection cloud microphysics gwd atmospheric gravity wave drag sso sub grid scale orographic effects mox methane oxidation and water vapor photolysis car Cariolle's linearized ozone chemistry
--

The time control for an atmospheric forcing by a process prc consists of three components, the time interval $dt_{-}prc$ for re-computing the forcing, and the start and end dates and times defining the interval $\lfloor sd_prc, ed_prc \rfloor$, in which the forcing is either computed, if the date/time coincides with the interval dt_prc, or recycled. Recycling means that the forcing stored from the last computation is used again. Outside of the interval the forcing is set to zero. If $dt_{-}prc$ is not specified, or an empty string or a string of blanks or an interval of length 0s, e.g. "PT0S" is given, then the forcing is switched off for the entire experiment and the start and end dates and times are irrelevant If sd-prc or ed-prc are not specified, or an empty string or a string of blanks are given, then the experiment start date and the experiment stop date are used, respectively.

Parameter	Type	Default	Unit	Description	Scope
mpi_phy_config(jg)%	C	""		This is the time interval in ISO 8601-2004	$run_nml/iforcing = 2$
dt_prc				format at which the forcing by the process prc	
				is computed.	
mpi_phy_config(jg)%	C	""		Defines the start date/time in ISO 8601-2004	$run_nml/iforcing = 2$
sd_prc				format of the interval (sd_prc, ed_prc) , in which	and $dt_{-}prc$; 0.000s
				the forcing by the process prc is computed in	
				intervals $dt_{-}prc$.	
mpi_phy_config(jg)%	C	""		Defines the end date/time in ISO 8601-2004	$run_nml/iforcing = 2$
ed_prc				format of the interval (sd_prc, ed_prc) , in which	and $dt_{-}prc$; 0.000s
				the forcing by the process prc is computed in	
				intervals dt - prc .	
mpi_phy_config(jg)%	ı	FALSE.		.TRUE. for sea-ice temperature calculation	$run_nml/iforcing = 2$
lice					
mpi_phy_config(jg)%	ı	FALSE.		TRUE. for mixed layer ocean	$run_nml/iforcing = 2$
lmlo					
mpi_phy_config(jg)%	Γ	FALSE.		.TRUE. for using the JSBACH land surface	$run_nml/iforcing = 2$
ljsb				model	
mpi_phy_config(jg)%	T	.FALSE.		TRUE. for AMIP boundary conditions	$run_nml/iforcing = 2$
lamip					

The namelist mpi_phy_nml is defined and read in: src/namelists/mo_mpi_phy_nml.f90

4.4.22 mpi_sso_nml

The parameterization of subgrid scale orographic (SSO) effects for the MPI physics is configured by a data structure $mpi_sso_config(jg=1:ndom)\%plarami_s$, which is a 1-dimensional array extending over all domains. The structure contains parameters providing control over the parametrized effects:

Parameter	Type	Default	Unit	Default Unit Description	Scope
mpi_sso_config(jg)%	R	40.	m	Minimum height difference of peak height and	mpi_phy_config(jg)%
gpicmea				mean height to activate the SSO	dt_{-sso} ; 0.000s
				parameterization.	
mpi_sso_config(jg)%	R	10.	m	Minimum standard deviation of the SSO height mpi_phy_config(jg)%	mpi_phy_config(jg)%
gstd				to activate the SSO parameterization.	dt_{-sso} ; 0.000s
mpi_sso_config(jg)%	R	0.05		Coefficient for orographic gravity wave drag.	mpi_phy_config(jg)%
gkdrag					$dt_{-sso} : 0.000s$
mpi_sso_config(jg)%	$_{ m R}$	0.		Coefficient for low level blocking.	$mpi-phy_config(jg)\%$
gkwake					dt_{-sso} ; 0.000s
mpi_sso_config(jg)%	$_{ m R}$	0.		Coefficient for low level lift.	$mpi-phy_config(jg)\%$
gklift					dt_{-sso} ; 0.000s

The namelist mpi_sso_nml is defined and read in: src/namelists/mo_mpi_sso_nml.f90

4.4.23 nonhydrostatic_nml (relevant if run_nml:iequations=3)

Parameter	Type	Default	Unit	Description	Scope
itime_scheme	I	4		Options for predictor-corrector time-stepping	
				scheme:	
				4: Contravariant vertical velocity is computed	iequations=3
				in the predictor step only, velocity tendencies	
				are computed in the corrector step only (most	
				efficient option)	
				5: Contravariant vertical velocity is computed	
				in both substeps (beneficial for numerical	
				stability in very-high resolution setups with	
				extremely steep slops, otherwise no significant	
				impact)	
				6: As 5, but velocity tendencies are also	
				computed in both substeps (no apparent	
				benefit, but more expensive)	

Parameter	Type	Default	Unit	Description	Scope
rayleigh-type	I	2		Type of Rayleigh damping 1: CLASSICAL (requires velocity reference state!)	
rayleigh_coeff	R(n_dom)	0.05		2: Klemp (2008) type Rayleigh damping coefficient $1/\tau_0$ (Klemp, Dudhia, Hassiotis: MWR136, pp.3987-4004);	
damp_height	R(n_dom)	45000	Ħ	finer resolution Height at which Rayleigh damping of vertical wind starts (needs to be adjusted to model top height; the damping layer should have a depth of at least 20 km when the model ton is above	
$\rm htop_moist_proc$	R	22500.0	m	the stratopause) Height above which moist physics and advection of cloud and precipitation variables	
hbot_qvsubstep	R	22500.0	ш	are turned off Height above which QV is advected with substepping scheme (must be at least as large	ihadv_tracer=22, 32, 42 or 52
vwind_offctr	R	0.15		as htop_moist_proc) Off-centering in vertical wind solver. Higher values may be needed for R2B5 or coarser grids	
rhotheta_offctr	R	-0.1		when the model top is above 50 km. Off-centering of density and potential temperature at interface level (may be set to	
veladv_offctr	R	0.25		0.0 for KZB6 or finer grids) Off-centering of velocity advection in corrector sten	
ivctype	П	2		Type of vertical coordinate: 1: Gal-Chen hybrid	
ndyn_substeps	I	ည		2: SLEVE (uses sleve_nml) number of dynamics substeps per fast-physics / transport step	

Parameter	Type	Default	Unit	Description	Scope
lhdiff_rcf	T	TRUE.		.TRUE.: Compute diffusion only at advection	
				time steps (in this case, divergence damping is	
				applied in the dynamical core)	
lextra_diffu	Γ	TRUE.		.TRUE.: Apply additional momentum diffusion	
				at grid points close to the stability limit for	
				vertical advection (becomes effective extremely	
				rarely in practice; this is mostly an emergency	
				fix for pathological cases with very large	
				orographic gravity waves)	
divdamp_fac	$_{ m R}$	0.0025		Scaling factor for divergence damping	$lhdiff_rcf = .TRUE.$
divdamp_order	I	4		Order of divergence damping:	$lhdiff_rcf = .TRUE.$
				2 = second-order divergence damping	
				4 = fourth-order divergence damping	
				24 = combined second-order and fourth-order	
				divergence damping and enhanced vertical	
				wind off-centering during the initial spinup	
				phase (does not allow checkpointing/restarting	
				earlier than 2.5 hours of integration)	
divdamp_type	I	3		Type of divergence damping:	$lhdiff_rcf = .TRUE.$
				2 = divergence damping acting on 2D	
				divergence	
				3 = divergence damping acting on 3D	
				divergence	
				32 = combination of 3D div. damping in the	
				troposphere with transition to 2D div. damping	
				in the stratosphere	
divdamp_trans_start	R	12500.		Lower bound of transition zone between 2D	$divdamp_type = 32$
				and 3D divergence damping	
divdamp_trans_end	${ m R}$	17500.		Upper bound of transition zone between 2D	$divdamp_type = 32$
				and 3D divergence damping	

Parameter	Type	Default	Unit	Description	Scope
nest_substeps	I	2		Number of dynamics substeps for the child	
				patches.	
				DO NOT CHANGE!!! The code will not work correctly with other values	
l-masscorr_nest	П	.FALSE.		TRUE: Apply mass conservation correction	
iadv_rhotheta	П	2		also in nested domain Advection method for rho and rhotheta:	
				1: simple second-order upwind-biased scheme	
				2: 2nd order Miura horizontal	
				3: 3rd order Miura horizontal (not	
				recommended)	
igradp_method	I	3		Discretization of horizontal pressure gradient:	
				1: conventional discretization with metric	
				correction term	
				2: Taylor-expansion-based reconstruction of	
				pressure (advantageous at very high resolution)	
				3: Similar discretization as option 2, but uses	
				hydrostatic approximation for downward	
				extrapolation over steep slopes	
				4: Cubic/quadratic polynomial interpolation	
				for pressure reconstruction	
				5: Same as 4, but hydrostatic approximation	
				for downward extrapolation over steep slopes	
l_zdiffu_t	L	TRUE.		.TRUE.: Compute Smagorinsky temperature	hdiff_order=3/5 .AND.
				diffusion truly horizontally over steep slopes	$lhdiff_temp = .true.$
thslp_zdiffu	R	0.025		Slope threshold above which truly horizontal	hdiff_order=3/5 .AND.
				temperature diffusion is activated	lhdiff_temp=.true.
					.AND. l_zdiffu_t=.true.
thhgtd_zdiffu	R	200	m	Threshold of height difference between	hdiff_order=3/5 .AND.
				neighboring grid points above which truly	lhdiff_temp=.true.
				horizontal temperature diffusion is activated	.AND. l_zdiffu_t=.true.
				(alternative criterion to thslp_zdiffu)	

Parameter	Type	Default Unit	Description	Scope
exner_expol	R	1./3.	Temporal extrapolation (fraction of dt) of	
			Exner function for computation of horizontal	
			pressure gradient. This damps horizontally	
			propagating sound waves. For R2B5 or coarser	
			grids, values between $1/2$ and $2/3$ are	
			recommended.	
l-open-ubc	Γ	.FALSE.	.TRUE.: Use open upper boundary condition	
			(rather than w=0) to allow vertical motions	
			related to diabatic heating to extend beyond	
			the model top	

Defined and used in: src/namelists/mo_nonhydrostatic_nml.f90

4.4.24 nwp_phy_nml

The switches for the physics schemes and the time steps can be set for each model domain individually. If only one value is specified, it is for the remaining model domains. If the time steps are not an integer multiple of the advective time step (dtime), then the time step of the copied to all child domains, implying that the same set of parameterizations and time steps is used in all domains. If the number of values given in the namelist is larger than 1 but less than the number of model domains, then the settings from the highest domain ID are used respective physics parameterization is automatically rounded to the next higher integer multiple of the advective time step.

Parameter	Type	Default Unit	Unit	Description	Scope
inwp-gscp	I (max.	1		cloud microphysics and precipitation	run_nml:iforcing =
	dom)			0: none	inwp
				1: hydci (COSMO-EU microphysics, 2-cat ice:	
				cloud ice, snow)	
				2: hydci_gr (COSMO-DE microphysics, 3-cat	
				ice: cloud ice, snow, graupel)	
				3: as 1, but with improved ice nucleation	
				scheme by C. Koehler	
				4: Two-moment microphysics by A. Seifert	
				9: Kessler scheme	

Parameter	Type	Default	Unit	Description	Scope
qi0	R	0.0	kg/kg	cloud ice threshold for autoconversion	inwp_gscp=1
dc0	R	0.0	kg/kg	cloud water threshold for autoconversion	inwp_gscp=1
mu_rain	$_{ m R}$	0.0		shape parameter in gamma distribution for rain	inwp_gscp>0
mu_snow	R	0.0		shape parameter in gamma distribution for	inwp_gscp>0
				Snow	
icpl_aero_gscp	I	0		0: off	currently only for
				1: simple coupling between autoconversion and	$inwp_gscp = 1$
				Tegen aerosol climatology; requires irad_aero=6	
				More advanced options are in preparation	
inwp_convection	I (max.	1		convection	run_nml:iforcing =
	dom)			0: none	inwp
				1: Tiedtke/Bechtold convection	
lshallowconv_only	L (max_dom)	.FALSE.		.TRUE.: use shallow convection only	inwp_convection = 1
ldetrain_conv_prec	Γ (max.	.FALSE.		.TRUE.: Activate detrainment of convective	$invp_convection = 1$
	dom)			rain and snow	
icapdcycl	Ι	0		Type of CAPE correction to improve diurnal	$invp_convection = 1$
				cycle for convection:	
				0 = none (IFS default prior to autumn 2013)	
				1 = intermediate testing option	
				2 = correctoins over land and water now	
				operational at ECMWF	
				3 = correction over land as in 2 restricted to	
				the tropics, no correction over water (this	
				choice optimizes the NWP skill scores)	
icpl_aero_conv	I	0		0: off	
				1: simple coupling between autoconversion and	
				Tegen aerosol climatology; requires irad-aero=6	
iprog_aero	I	0		0: off	
				1: simple prognostic aerosol scheme, based on	
				2D aerosol optical depth fields of Tegen	
	_		_	cumatology, requires man-aero—o	_

Parameter	Type	Default	Unit	Description	Scope
icpl_o3_tp	I	1		0: off	$irad_03 = 7 \text{ or } 9$
				1: simple coupling between the ozone mixing ratio and the thermal tropopause, restricted to the extratropics	
inwp_cldcover	I (max.	П		cloud cover scheme for radiation 0. no clouds (only OV)	run_nml:iforcing =
	aouit)			1: diagnostic cloud cover (by Martin Koehler)	Jam
				2: prognostic total water variance (not yet	
				statisty)	
				4: clouds as in turbulence (turbdiff)	
				5: grid scale clouds	
inwp_radiation	I (max_	1		radiation	run_nml:iforcing =
	dom)			0: none	inwp
				1: RRTM radiation	
				2: Ritter-Geleyn radiation	
				3: PSRAD radiation	
inwp_satad	Ι			saturation adjustment	run_nml:iforcing =
				0: none	inwp
				1: saturation adjustment at constant density	
$ \text{inwp_turb} $	I (max_	1		vertical diffusion and transfer	run_nml:iforcing =
	dom)			0: none	inwp
				1: COSMO diffusion and transfer	
				2: GME turbulence scheme	
				3: EDMF-DUALM (work in progress)	
				5: Classical Smagorinsky diffusion	
inwp_sso	I (max_	1		subgrid scale orographic drag	run_nml:iforcing =
	dom)			0: none	inwp
				1: Lott and Miller scheme (COSMO)	
inwp_gwd	I (max-	1		non-orographic gravity wave drag	run_nml:iforcing =
	dom)			0: none	inwp
				1: Orr-Ern-Becntold-scheme (IFS)	

Parameter	Type	Default	Unit	Description	Scone
	7 J P.C	ormano,			
inwp_surface	I (max_	-		surface scheme	$run_nml:iforcing =$
	dom)			0: none	inwp
				1: TERRA	
ustart_raylfric	R	160.0	$\mathrm{m/s}$	wind speed at which extra Rayleigh friction	$\operatorname{inwp_gwd} > 0$
				starts	
efdt_min_raylfric	R	10800.	∞	minimum e-folding time of Rayleigh friction	$\operatorname{inwp_gwd} > 0$
				(effective for $u > ustart_raylfric + 90 m/s$)	
latm_above_top	L (max_	FALSE.		.TRUE.: take into account atmosphere above	inwp_radiation > 0
	dom)			model top for radiation computation	
itype_z0	Ι	2		Type of roughness length data used for	$inwp_turb > 0$
				turbulence scheme:	
				1 = land-cover-related roughness including	
				contribution from sub-scale orography (does	
				not account for tiles)	
				2 = land-cover-related roughness based on	
				tile-specific landuse class	
				3 = land-cover-related roughness based on	
				tile-specific landuse class including contribution	
				from sub-scale orography	
dt_{-conv}	R (max_	.009	∞	time interval of convection call	run_nml:iforcing =
	dom)			currently each subdomain has the same value	inwp
dt_rad	R (max_	1800.	∞	time interval of radiation call	run_nml:iforcing =
	dom)			currently each subdomain has the same value	inwp
$ m dt_sso$	R (max	1200.	∞	time interval of sso call	run_nml:iforcing =
	dom)			currently each subdomain has the same value	inwp
$\mathrm{dt_gwd}$	m R~(max	1200.	w	time interval of gwd call	run_nml:iforcing =
	dom)			currently each subdomain has the same value	inwp
lrtm_filename	C(:)	m ``rrtmg-		NetCDF file containing longwave absorption	
		lw.nc"		coefficients and other data for RRTMG_LW	
				k-distribution model.	
cldopt_filename	C(:)	"ECHAM		NetCDF file with RRTM Cloud Optical	
		6-CldOpt		Properties for ECHAM6.	
		Props.nc"			

Defined and used in: src/namelists/mo_nwp_phy_nml.f90

4.4.25 nwp_tuning_nml

Please note: These tuning parameters are NOT domain specific.

Parameter	Type	Default	Unit	Description	Scope
SSO (Lott and Miller)					
tune_gkwake	R	1.5		low level wake drag constant	run_nml:iforcing = inwp
tune-gkdrag	Я	0.075		gravity wave drag constant	run_nml:iforcing = inwp
GWD (Warner McIntyre)	e)				
tune-gfluxlaun	R	2.50e-3		total launch momentum flux in each azimuth	run_nml:iforcing =
				(110-0 A 1 -0)	Awiii
Grid scale microphysics (one moment	cs (one mo	ment)			
tune_zceff_min	R	0.075		Minimum value for sticking efficiency	run_nml:iforcing = inwp
tune_v0snow	R	25.0		factor in the terminal velocity for snow	run_nml:iforcing = inwp
tune_zvz0i	R	1.25	s/m	Terminal fall velocity of ice	run_nml:iforcing =
					inwp
Convection scheme					
tune_entrorg	R	1.85e-3	$1/\mathrm{m}$	Entrainment parameter valid for dx=20 km	run_nml:iforcing =
				(depends on model resolution)	inwp
tune_capdcfac_et	m R	0.125		Fraction of CAPE diurnal cycle correction	icapdcycl = 3
				applied ill tile extractopies	
tune_rhebc_land	R	0.75		RH threshold for onset of evaporation below	$run_nml:iforcing =$
				cloud base over land	inwp
tune_rhebc_land_trop	R	0.70		RH threshold for onset of evaporation below	run_nml:iforcing =
				cloud base over land in the tropics	inwp

Parameter	$\mid { m Type}$	Default	Unit	Description	Scope
tune_rhebc_ocean	R	0.85		RH threshold for onset of evaporation below	run_nml:iforcing =
				cloud base over sea	inwp
tune_rhebc_ocean_trop	R	08.0		RH threshold for onset of evaporation below	run_nml:iforcing =
				cloud base over sea in the tropics	inwp
tune_rcucov	R	0.05		Convective area fraction used for computing	run_nml:iforcing =
				evaporation below cloud base	inwp
tune_rcucov_trop	R	0.05		Convective area fraction used for computing	run_nml:iforcing =
				evaporation below cloud base in the tropics	inwp
tune_texc	R	0.125	K	Excess value for temperature used in test	run_nml:iforcing =
				parcel ascent	inwp
tune_qexc	R	0.0125		Excess fraction of grid-scale QV used in test	run_nml:iforcing =
				parcel ascent	inwp
Misc					
itune_albedo	I	0		MODIS albedo tuning	run_nml:iforcing =
				0: None	inwp
				1: dimmed sahara	albedo_type=2
tune_minsnowfrac	R	0.125		Minimum value to which the snow cover	lnd_nml:idiag_snowfrac
				fraction is artificially reduced in case of melting	= 20/30/40
				show	
IAU					
max_freshsnow_inc	R	0.025		Maximum allowed freshsnow increment per	init_mode=5
				analysis cycle (positive or negative)	(MODE_IAU)

Defined and used in: src/namelists/mo_nwp_tuning_nml.f90

4.4.26 output_nml (relevant if run_nml/output='nml')

Please note: There may be several instances of output_nml in the namelist file, every one defining a list of variables with separate attributes for output.

Parameter	Type	Default	Unit	Description S	Scope
dom	I(:)	-1		Array of domains for which this name-list is	
				used. If not specified (or specified as -1 as the first array member), this name-list will be used	
				for all domains.	
				Attention: Depending on the setting of the	
				parameter l_output_phys_patch these are either	
		,		logical or physical domain numbers!	
file_interval	C	\r \r		Defines the length of a file in terms of an	
				ISO-8601 duration string. An example for this	
				time stamp format is given below. This	
				namelist parameter can be set instead of	
				steps_per_file.	
filename_format	C	see		Output filename format. Includes keywords	
		descrip-		path, output_filename, physdom, etc. (see	
		tion.		below). Default is	
				<pre><output_filename>_DOM<physdom>_<levtype>_</levtype></physdom></output_filename></pre>	
				<jfile></jfile>	
filename_extn	C	"default"		User-specified filename extension (empty string	
				also possible). If this namelist parameter is	
				chosen as "default", then we have ".nc" for	
				NetCDF output files, and ".grb" for GRIB1/2.	
filetype	Ι	4		One of CDI's FILETYPE_XXX constants.	
				Possible values:	
				2=FILETYPE_GRB2,	
				4=FILETYPE_NC2,	
				5=FILETYPE_NC4	

Parameter	Type	Default	Unit	Description	Scope
m levels	D	None		Model level indices (optional). Allowed is a comma- (or semicolon-) separated list of integers, and of integer ranges like "1020". One may also use the keyword "nlev" to denote the maximum integer (or, equivalently, "n" or "N"). Furthermore, arithmetic expressions like "(nlev - 2)" are possible.	
				Basic example: m_levels = "1,3,510,20(nlev-2)"	
h_levels	$\mathbf{R}(:)$	None	ш	height levels	
p_levels	R(:)	None	Pa	pressure levels	
i_levels	R(:)	None	X	isentropic levels	
ml_varlist	Q(:)	None		Name of model level fields to be output.	
varlist varlist	(i) (ii)	None None		Name of neight level fields to be output. Name of pressure level fields to be output.	
il_varlist	C(:)	None		Name of isentropic level fields to be output.	
include_last	ı	TRUE.		Flag whether to include the last time step	
mode	П	2		1 = forecast mode, $2 = $ climate mode In climate mode the time axis of the output file	
				is set to TAXIS_ABSOLUTE. In forecast mode it is set to TAXIS_RELATIVE. Till now the	
				forecast mode only works if the output is at multiples of 1 hour	

Parameter	Type	Default	Unit	Description	Scope
taxis_tunit	I	2		IS_RELATIVE time axis.	mode=1
				$1 = TUNIT_SECOND$	
				$2 = \text{TUNIT_MINUTE}$	
				$5 = TUNIT_HOUR$	
				$9 = TUNIT_DAY$	
				For a complete list of possible values see	
				cdilib.c	
$output_bounds$	R(k*3)	None		Post-processing times: start, end, increment.	
				We choose the advection time step matching or	
				following the requested output time, therefore	
				we require output_bounds(3) > dtime.	
				Multiple triples are possible in order to define	
				multiple starts/ends/intervals. See namelist	
				parameters output_start, output_end,	
				output_interval for an alternative	
				specification of output events.	
output_time_unit	Ι			Units of output bounds specification.	
				1 = second	
				2 = minute	
				3 = hour	
				4 = dav	
				$5 \equiv month$	
output_filename	C	None		Output filename prefix (which may include	
ı				path). Domain number, level type, file number	
				and extension will be added, according to the	
				format given in namelist parameter	
				"filename_format".	
output_grid	Τ	.FALSE.		Flag whether grid information is added to	
				output.	

Parameter	Type	Default	Unit	Description	Scope
output_start	C(:)	; ;		ISO8601 time stamp for begin of output. An	
				example for this time stamp format is given	
				below. More than one value is possible in order	
				to define multiple start/end/interval triples.	
				See namelist parameter output_bounds for an	
				alternative specification of output events.	
output_end	C(:)	,,,		ISO8601 time stamp for end of output. An	
				example for this time stamp format is given	
				below. More than one value is possible in order	
				to define multiple start/end/interval triples.	
				See namelist parameter output_bounds for an	
				alternative specification of output events.	
output_interval	C(:)	,,,		ISO8601 time stamp for repeating output	
				intervals. We choose the advection time step	
				matching or following the requested output	
				time, therefore we require output_bounds(3)	
				> dtime. An example for this time stamp	
				format is given below. More than one value is	
				possible in order to define multiple	
				start/end/interval triples. See namelist	
				parameter output_bounds for an alternative	
				specification of output events.	
operation	O	None		Choose "mean" for generating time averaged	
				output for the given list of variables or groups.	
				The corresponding interval is the	
				output_interval. Supported are 2D and 3D	
				fields on model levels of the atmosphere and	
				land model. Any other value than mean will be	
				ignored.	

Parameter	Type	Default	Unit	Description 5	Scope
pe_placement_il	I(:)	-1		Advanced output option: Explicit assignment	
				of output MPI ranks to the isentropic level	
				output file. At most stream_partitions_il	
				different ranks can be specified. See namelist	
				parameter pe_placement_ml for further details.	
pe_placement_hl	I(:)	-1		Advanced output option: Explicit assignment	
				of output MPI ranks to the height level output	
				file. At most stream_partitions_hl different	
				ranks can be specified. See namelist parameter	
				pe_placement_ml for further details.	
pe_placement_ml	I(:)	-1		Advanced output option: Explicit assignment	
				of output MPI ranks to the model level output	
				file. At most stream_partitions_ml different	
				ranks can be specified, out of the following list:	
				0 (num_io_procs - 1). If this namelist	
				parameters is not provided, then the output	
				ranks are chosen in a Round-Robin fashion	
				among those ranks that are not occupied by	
				explicitly placed output files.	
pe_placement_pl	I(:)	-		Advanced output option: Explicit assignment	
				of output MPI ranks to the pressure level	
				output file. At most stream_partitions_pl	
				different ranks can be specified. See namelist	
				parameter pe_placement_ml for further details.	

Parameter	Type	Default	Unit	Description	Scope
ready_file	C	'default'		A ready file is a technique for handling	
				dependencies between the NWP processes. The	
				completion of the write process is signalled by	
				creating a small file with name ready_file.	
				Different output_nml's may be joined together	
				to form a single ready file event. The setting of	
				ready_file = "default" does not create a	
				ready file. The ready file name may contain	
				string tokens <path>, <datetime>, <ddhhmmss></ddhhmmss></datetime></path>	
				which are substituted as described for the	
				namelist parameter filename_format.	
reg_def_mode	I	0		Specify if the "delta" value prescribes an	remap=1
				interval size or the total *number* of intervals:	
				0: switch automatically between increment and	
				no. of grid points, 1: reg_lon/lat_def(2)	
				specifies increment, 2: reg_lon/lat_def(2)	
				specifies no. of grid points.	
remap	I	0		interpolate horizontally	
				0: none	
				1: to regular lat-lon grid	
north_pole	R(2)	0,90		definition of north pole for rotated lon-lat grids	
				([longitude, latitude].	
$ m reg_lat_def$	R(3)	None		start, increment, end latitude in degrees.	remap=1
				Alternatively, the user may set the number of	
				grid points instead of an increment. Details for	
				the setting of regular grids is given below	
				together with an example.	
reg_lon_def	R(3)	None		The regular grid points are specified by three	remap=1
				values: start, increment, end given in degrees.	
				Alternatively, the user may set the number of	
				grid points instead of an increment. Details for	
				the setting of regular grids is given below	
				together with an example.	

Parameter	Type	Default	Unit	Description	Scope	
steps_per_file	I	-1		Max number of output steps in one output file.	•	
				If this number is reached, a new output file will		
				be opened.		
steps_per_file_inclfirst	L	see		Defines if first step is counted wrt.		
		descr.		steps_per_file files count. The default is		
				.FALSE. for GRIB2 output, and .TRUE.		
				otherwise.		
stream_partitions_hl	Ι	1		Splits height level output of this namelist into		
				several concurrent alternating files. See		
				namelist parameter stream_partitions_ml for		
				details.		
stream_partitions_il	I	1		Splits isentropic level output of this namelist		
				into several concurrent alternating files. See		
				namelist parameter stream partitions_ml for		
				details.		
stream_partitions_ml	Ι	1		Splits model level output of this namelist into		
				several concurrent alternating files. The output		
				is split into N files, where the start date of		
				part i gets an offset of		
				$(i-1)*$ output_interval. The output interval		
				is then replaced by $N * \text{output_interval}$, the		
				include_last flag is set to .FALSE., the		
				steps_per_file_inclfirst flag is set to		
				.FALSE., and the steps_per_file counter is		
				set to 1.		
stream_partitions_pl	I	1		Splits pressure level output of this namelist		
				into several concurrent alternating files. See		
				namelist parameter stream_partitions_ml for		
				details.		
rbf_scale	R	-1.		Explicit setting of RBF shape parameter for	$interpol_nml:rbf_scale_mode_ll=$	de_ll=
				interpolated lon-lat output. This namelist		
				parameter is only active in combination with		
				interpol_nml:rbt_scale_mode_ll=3.		

Defined and used in: src/io/shared/mo_name_list_output_init.f90

Horizontal interpolation onto regular grids is possible through the namelist setting remap=1, where the Interpolation onto regular grids: mesh is defined by the parameters

- reg_lon_def: mesh latitudes in degrees,
- reg_lat_def: mesh longitudes in degrees,
- north_pole: definition of north pole for rotated lon-lat grids.

The regular grid points in reg_lon_def, reg_lat_def are each specified by three values, given in degrees: start, increment, end. The mesh then contains all grid points $start + k * increment \le end$, where k is an integer. Instead of defining an increment it is also possible to prescribe the number of grid points.

- Setting the namelist parameter reg_def_mode=0: Switch automatically from increment specification to no. of grid points, when the reg_lon/lat_def(2) value is larger than 5.0.
- 1: reg_lon/lat_def(2) specifies increment
- 2: reg_lon/lat_def(2) specifies no. of grid points

For longitude values the last grid point is omitted if the end point matches the start point, e.g. for 0 and 360 degrees.

Examples

ocal grid with 0.5 degree increment:

global grid with 720x361 grid points:

reg_lon_def = 0.,720,360. reg_lat_def = -90.,360,90.

reg_lon_def = -30.,0.5,30. reg_lat_def = 90.,-0.5, -90. Time stamp format: The namelist parameters output_start, output_end, output_interval allow the specification of time stamps according to ISO 8601. The general format for time stamps is YYYY-MM-DDThh:mm:ss where Y: year, M: month, D: day for dates, and hh: hour, mm: minute, ss: second for time strings. The general format for durations is PnYnMnDTnHnMnS. See, for example, http://en.wikipedia.org/wiki/ISO_8601 for details and further specifications.

NOTE: as the mtime library underlaying the output driver currently has some restrictions concerning the specification of durations:

- 1. Any number n in PnYnMnDTnHnMnS must have two digits. For instance use "PT06H" instead of "PT6H"
- In a duration string PnyearYnmonMndayDTnhrHnminMnsecS the numbers nxyz must not pass the carry over number to the next larger time unit: 0;=nmon;=12, 0;=nhr;=23, 0;=nmin;=59, 0;=nsec;=59.999. For instance use "P01D" instead of "PT24H", or "PT01M" instead "S09Tq" fo

Soon the formatting problem will be resolved and the valid number ranges will be enlarged. (2013-12-16).

Examples

2013-10-27T13:41:00Z POODTO6HOOMOOS date and time representation (output_start, output_end) duration (output_interval)

Variable Groups

Keyword "group:": Using the "group:" keyword for the namelist parameters ml_varlist, hl_varlist, pl_varlist, sets of common variables can be added to the output

output of all variables (caution: do not combine with mixed vertical interpolation) basic atmospheric variables on model levels group:atmo_pl_vars group:atmo_ml_vars group:all

additional prognostic variables of the nonhydrostatic model same set as atmo_ml_vars, but expect height same set as atmo-ml-vars, but except pres

derived atmospheric variables group:atmo_derived_vars

group:nh_prog_vars

group:atmo_zl_vars

snow variables group:phys_tendencies group:cloud_diag group:land_vars group:pbl_vars

group:multisnow_vars group:snow_vars

group:additional_precip_vars group:dwd_fg_atm_vars

DWD first guess fields (atmosphere) multi-layer snow variables

group:precip_vars

group:rad_vars

OWD first guess fields (surface/soil) ART radioactive tracer fields ART volcanic ash fields group:dwd_fg_sfc_vars group: ART_AERO_RADIO group:ART_AERO_VOLC

ART mineral dust aerosol fields ART sea salt aerosol fields group:prog_timemean group: ART_AERO_SEAS group:ART_AERO_DUST

time mean variables from prog_timemean,tracer_timemean, echam_timemean group:tracer_timemean group:echam_timemean group:atmo_timemean

sime mean output: most echam surface variables sime mean output: temp, u, v, rho time mean output: qv, qc, qi

"tiles:t_g" (read: "tiles of t_g") automatically adds all t_g_t_X fields to the output. Here, X is a placeholder for the Keyword "tiles:": The "tiles:" keyword allows to add all tiles of a specific variable to the output, without the need to specify all tile tile number. Make sure to specify the name of the aggregated variable rather than the name of the corresponding tile container (i.e. in the given example it must be t_g , and not $t_g(t)$. fields separately. E.g.

There exists a special syntax which allows to remove variables from the output list, e.g. if these undesired variables were contained in a previously selected group.

Typing "-jvarname;" (for example "-temp") removes the variable from the union set of group variables and other selected variables Note that typos are not detected but that the corresponding variable is simply not removed!

Keyword substitution in output filename (filename_format):

substituted by level type "ML", "PL", "HL", "IL" substituted by output file counter substituted by output_filename substituted by physical patch ID substituted by model_base_dir like levtype, but in lower case output_filename levtype_l levtype physdom file

substituted by ISO-8601 date-time stamp in format YYYY-MM-DDThh:mm:ss.ssz substituted by ISO-8601 date-time stamp in format YYYYMMDDThhmmssZ

datetime2

datetime

substituted by ISO-8601 date-time stamp in format YYYYMMDDThhmmss.sssZ substituted by relative day-hour-minute-second string substituted by relative hour-minute-second string

If namelist is split into concurrent files: number of stream partitions.

If namelist is split into concurrent files: stream partition index of this file. If namelist is split into concurrent files: substituted by the file counter

(like in jfile), which an "unsplit" namelist would have produced

ifile_partition npartitions total_index datetime3 ddhhmmss hhhmmss

4.4.27 parallel_nml

Parameter	Type	Default	Unit	Description	Scope
nproma	П	1		chunk length	
n_ghost_rows	Ι	1		number of halo cell rows	
division_method	Ι	1		method of domain decomposition	
				0: read in from file	
				1: use built-in geometric subdivision	
division_file_name	C			Name of division file	$division_method = 0$
ldiv_phys_dom	П	.TRUE.		.TRUE.: split into physical domains before	$division_method = 1$
				computing domain decomposition (in case of	
				merged domains)	
				(This reduces load imbalance; turning off this	
				option is not recommended except for very	
				small processor numbers)	
p_test_run	J	.FALSE.		TRUE. means verification run for MPI	
				parallelization (PE 0 processes full domain)	
$1_{ m test_openmp}$	T	.FALSE.		if .TRUE. is combined with	$p_{-test_run} = .TRUE.$
				p_test_run=.TRUE. and OpenMP	
				parallelization, the test PE gets only 1 thread	
				in order to verify the OpenMP parallelization	
l_log_checks	T	.FALSE.		if .TRUE. messages are generated during each	
				synchonization step (use for debugging only)	
$1_{ m fast_sum}$	П	.FALSE.		if .TRUE., use fast (not	
				processor-configuration-invariant) global	
				summation	
use_dycore_barrier	П	.FALSE.		if .TRUE., set an MPI barrier at the beginning	
				of the nonhydrostatic solver (do not use for	
				production runs!)	
itype_exch_barrier	П	0		1: set an MPI barrier at the beginning of each	
				MPI exchange call	
				2: set an MPI barrier after each MPI WAIT call	
				3: 1+2 (do not use for production runs!)	_

Parameter	Type	Default	Unit	Description	Scope
iorder_sendrecv	П	П		Sequence of send/receive calls:	
				1 = irecv/send	
				2 = isend/recv	
				3 = isend/irecv	
itype_comm	I	1		1: use local memory for exchange buffers	
				3: asynchronous halo communication for	
				dynamical core (currently deactivated)	
num_io_procs	I	0		Number of I/O processors (running exclusively	
				for doing I/O)	
${\tt num_restart_procs}$	I	0		Number of restart processors (running	
				exclusively for doing restart)	
num_prefetch_proc	I	0		Number of processors for prefetching of	itype_latbc ≥ 1
				boundary data asynchronously for a limited	
				area run (running exclusively for reading Input	
				boundary data. Maximum no of processors	
				used for it is limited to 1).	
pio-type	I	1		Type of parallel I/O. Only used if number of	
				I/O processors greater than number of	
				domains. Experimental!	
use_icon_comm	П	.FALSE.		Enable the use of MPI bulk communication	
				through the icon_comm_lib	
icon_comm_debug	Г	FALSE.		Enable debug mode for the icon_comm_lib	
max_send_recv-	Ι	131072		Size of the send/receive buffers for the	
_buffer_size				icon_comm_lib.	
use_dp_mpi2io	П	.FALSE.		Enable this flag if output fields shall be	
				gathered by the output processes in DOUBLE	
				PRECISION.	
restart_chunk_size	I	П		(Advanced namelist parameter:) Number of	
				levels to be buffered by the asynchronous	
				restart process. The (asynchronous) restart is	
				capable of writing and communicating more	
				than one 2D slice at once.	

Defined and used in: src/namelists/mo_parallel_nml.f90

Scope tion of the random seed from column to column completely randomly in order to reduce errors switch for diagnostics of aerosol forcing in the chosen g–points per column and radiation call integer number that influences the perturbanumber of g-points in Monte-Carlo spectral solar spectral range (lradforcing(1)) and the number of g-points in Monte-Carlo spectral lw.spec.samp = 2: Monte-Carlo spec- tral integration (MSCI); lw-gpts_ts randomly $lw_spec_samp = 1$: standard broad band thermal spectral range (lradforcing(2)). sampling of spectral bands in radiation $lw_spec_samp = 3$: choose g-points not integration for thermal radiation, see integration for solar radiation, see calculation for thermal radiation in the surface radiative fluxes lw_spec_samp sw_spec_samp Description sampling Unit FALSE. Default 0 Type $\Gamma(2)$ lw_spec_samp Parameter lradforcing $sw-gpts_ts$ lw_-gpts_-ts rad_perm

4.4.28 psrad_nml

9	
Scope	
Default Unit Description	sampling of spectral bands in radiation calculation for solar radiation sw_spec_samp = 1: standard broad band sampling sw_spec_samp = 2: Monte-Carlo spectral integration (MSCI); lw_gpts_ts randomly chosen g_points per column and radiation call sw_spec_samp = 3: choose g_points not completely randomly in order to reduce errors in the surface radiative fluxes
Unit	
Default	⊢
Type	П
Parameter	sw_spec_samp

Defined and used in: src/echam_phy_psrad/mo_psrad_radiation.f90

4.4.29 psrad_orbit_nml

Scope			orbit of		orbit is		letric	liation	···
Default Unit Description	eccentricity of earth's orbit	obliquity of earth in degrees	switch on (.TRUE.) the (real) observed orbit of	the earth (not idealized) or switch it off	(.FALSE.). In the latter case, a Kepler orbit is	used.	switch on (.TRUE.) a spherically symmetric	irradiation from all sides or use an irradiation	by a point source like the sun (.FALSE.).
Unit									
Default	0.016715	23.44100	.TRUE.				FALSE.		
$\mid \text{Type}$	R	R	L				ı		
Parameter	cecc	cobld	l_orbvsop87				l_sph_symm_irr		

Defined and used in: src/namelists/mo_psrad_radiation.f90

4.4.30 radiation_nml

Parameter	Type	Default	Unit	Description	Scope
ldiur	L	.TRUE.		switch for solar irradiation:	
				.TRUE.:diurnal cycle,	
				.FALSE.:zonally averaged irradiation	
nmonth	I	0		0: Earth circles on orbit	
				1-12: Earth orbit position fixed for specified	
				month	
lyr-perp	Г	FALSE.		.FALSE.: transient Earth orbit following	
				VSOP87	
				.TRUE.: Earth orbit of year yr-perp of the	
				VSOP87 orbit is perpertuated	
yr-perp	L	66666-		year used for $lyr_perp = .TRUE$.	
isolrad	I	0		Insolation scheme	
				0: Use original SRTM insolation.	
				1: Use insolation from external file containing	
				the spectrally resolved insolation (monthly	
				means)	
				2: Use preindustrial insolation as in CMIP5	
				(average from $1844-1856$)	
				3: Use insolation for AMIP-type CMIP5	
				simulation (average from 1979–1988)	
				4: Use insolation for RCE-type simulation with	
				$\cos(\text{zenith angle}) = \text{pi}/4 \text{ (with PSRAD)}$: use	
				"4" if the diurnal cycle is switched on)	
				5: Use insolation for RCE-type simulation with	
				PSRAD if the diurnal cycle is switched off.	

Parameter	Type	Default	Unit	Description	Scope
izenith	Н	4		Choice of zenith angle formula for the radiative transfer computation. 0: Sun in zenith everywhere 1: Zenith angle depends only on latitude 2: Zenith angle depends only on latitude. Local time of day fixed at 07:14:15 for radiative transfer computation (sin(time of day) = 1/pi 3: Zenith angle changing with latitude and time of day 4: Zenith angle and irradiance changing with season, latitude, and time of day (iforcing=inwp only)	
albedo_type	Н	1		Type of surface albedo 1: based on soil type specific tabulated values (dry soil) 2: MODIS albedo	iforcing=inwp
direct_albedo	н	4		Direct beam surface albedo. Options mainly differ in terms of their solar zenith angle (SZA) dependency) 1: SZA dependency following Ritter-Geleyn; applied to unconditionally all grid points 2: SZA dependency following Zaengl (pers. comm.). Same as 1 for water, but for 'rough surfaces' over land the direct albedo is not allowed to exceed the corresponding broadband diffuse albedo. 3: SZA dependency following Yang (2008) for snow-free land points. Same as 1 for water/ice and 2 for snow. 4: SZA dependency following Briegleb (1992) for snow-free land points. Same as 1 for water/ice and 2 for snow.	iforcing=inwp albedo_type=2

Parameter	Type	Default	Unit	ult Unit Description	Scope
icld_overlap	I	2		Method for cloud overlap calculation in	iforcing=inwp
				shortwave part of RRTM	inwp_radiation=1
				1: maximum-random overlap	
				2: generalized overlap (Hogan, Illingworth,	
				2000)	
				3: maximum overlap	
				4: random overlap	

1 Switches for the concentration of radiative 2 agents 3 irad_Ayz = 0: set to zero 3 dirad_Ab2 = 1: vapor, cloud water and cloud ice 6 from tracer variables 2 concentration given by 2 concentration given by 2 vmr_co2/ch4/n2o/o2/cfc11/cfc12 = 2: 2 concentration given by vmr_ch4/n2o 2 irad_co2/cfc11/cfc12 = 4: 2 irad_co2/cfc11/cfc12 = 4: time dependent 2 concentration from greenhouse gas file 3 irad_co4/n2o = 4: time dependent tanh-profile 3 with surface concentration from greenhouse gas 4 irad_co3 = 2: ozone climatology from MPI 3 irad_co3 = 2: ozone climatology from MPI 3 geographical distribution and Fourier series for 3 seasonal cycle for run_nnl/flocing = 3 (NWP) 3 irad_co3 = 7: GEMS ozone climatology from 3 irad_co3 = 7: GEMS ozone climatology from 3 irad_co3 = 7: GEMS ozone climatology from 3 irad_co3 = 9: MACC ozone climatology from 4 irad_co3 = 9: MACC ozone climatology from 5 irad_co3 = 10: Linearized ozone clemistry (ART 6 extension necessary) for run_nnl/iforcing = 3 6 irad_cos_nnecessary) for run_nnl/iforcing =	туре	Default	Unit	Description	Scope
	I	1		Switches for the concentration of radiative	
		2		agents	
		3		$irad_xyz = 0$: set to zero	
		3		irad_h2o = 1: vapor, cloud water and cloud ice	
		0		from tracer variables	
		2		$irad_co2 = 1$: CO_2 from tracer variable	
		2		$irad_{co2}/ch4/n2o/o2/cfc11/cfc12 = 2$:	
vmr.co2/ch4/n2o/o2/cfc11/cfc12 irad_ch4/n2o = 3: tanh-profile with surface concentration given by vmr.ch4/n2o irad_co2/cfc11/cfc12 = 4: time dependent concentration from greenhouse gas file irad_ch4/n2o = 4: time dependent tanh-profile with surface concentration from greenhouse gas file irad_o3 = 2: ozone climatology from MPI irad_o3 = 4: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_o3 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3		2		concentration given by	
irad_co4/n2o = 3: tanh-profile with surface concentration given by vmr_ch4/n2o irad_co2/cfc11/cfc12 = 4: time dependent concentration from greenhouse gas file irad_ch4/n2o = 4: time dependent tanh-profile with surface concentration from greenhouse gas file irad_o3 = 2: ozone climatology from MPI irad_o3 = 4: ozone climatology from MPI irad_o3 = 6: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_o3 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				$vmr_co2/ch4/n2o/o2/cfc11/cfc12$	
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irad_co2/cfc11/cfc12 = 4: time dependent concentration from greenhouse gas file irad_ch4/n2o = 4: time dependent tanh-profile with surface concentration from greenhouse gas file irad_o3 = 2: ozone climatology from MPI irad_o3 = 4: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_o3 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 8: ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				concentration given by vmr_ch4/n2o	
concentration from greenhouse gas file irad_ch4/n2o = 4: time dependent tanh-profile with surface concentration from greenhouse gas file irad_o3 = 2: ozone climatology from MPI irad_o3 = 4: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_o3 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 8: ozone climatology for AMIP irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				$irad_{co2}/cfc11/cfc12 = 4$: time dependent	
irad_ch4/n2o = 4: time dependent tanh-profile with surface concentration from greenhouse gas file irad_o3 = 2: ozone climatology from MPI irad_o3 = 4: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_o3 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 8: ozone climatology for AMIP irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				concentration from greenhouse gas file	
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file irad_03 = 2: ozone climatology from MPI irad_03 = 4: ozone clim for Aqua Planet Exp irad_03 = 6: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_03 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 8: ozone climatology for AMIP irad_03 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				with surface concentration from greenhouse gas	
irad_o3 = 4: ozone clim for Aqua Planet Exp irad_o3 = 6: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_o3 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 8: ozone climatology for AMIP irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				file irad_o3 = 2: ozone climatology from MPI	
irad-03 = 6: ozone climatology with T5 geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad-03 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad-03 = 8: ozone climatology for AMIP irad-03 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad-03 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad-03 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				$irad_03 = 4$: ozone clim for Aqua Planet Exp	
geographical distribution and Fourier series for seasonal cycle for run_nml/iforcing = 3 (NWP) irad_03 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 8: ozone climatology for AMIP irad_03 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				$irad_03 = 6$: ozone climatology with T5	
seasonal cycle for run_nml/iforcing = 3 (NWP) irad_03 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				geographical distribution and Fourier series for	
irad_o3 = 7: GEMS ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 8: ozone climatology for AMIP irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				seasonal cycle for run_nml/iforcing = 3 (NWP)	
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irad_o3 = 8: ozone climatology for AMIP irad_o3 = 9: MACC ozone climatology (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				IFS) for run nml/iforcing = 3 (NWP)	
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IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_03 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				$irad_03 = 9$: MACC ozone climatology (from	
irad_o3 = 79: Blending between GEMS and MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				IFS) for run nml/iforcing = 3 (NWP)	
MACC ozone climatologies (from IFS) for run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				$irad_03 = 79$: Blending between GEMS and	
run_nml/iforcing = 3 (NWP) irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = 3				MACC ozone climatologies (from IFS) for	
irad_o3 = 10: Linearized ozone chemistry (ART extension necessary) for run_nml/iforcing = $\frac{1}{3}$				$run_nml/iforcing = 3 (NWP)$	
extension necessary) for run nml/iforcing = 3				$irad_03 = 10$: Linearized ozone chemistry (ART	
				extension necessary) for run_nml/iforcing = 3	

Parameter	Type	Default	Unit	Description	Scope
vmr_co2 vmr_ch4 vmr_n2o	Я	348.0e-6 1650.0e-9 306.0e-9		Volume mixing ratio of the radiative agents	
vmr_o2 vmr_cfc11 vmr_cfc12		0.20946 214.5e-12 371.1e-12			
fh2o fco2 fch4 fn.2c	ಜ			Scaling factors for concentrations used in radiation	run_nml/iforcing=2 (ECHAM)
fo3 fo2 fcfc					
irad_aero	H	67		Aerosols 1: prognostic variable 2: global constant 3: externally specified 5: Tanre aerosol climatology for run_nml/iforcing = 3 (NWP) 6: Tegen aerosol climatology for run_nml/iforcing = 3 (NWP) .AND. itopo = 1 9: ART online aerosol radiation interaction, uses Tegen for aerosols not chosen to be represented in ART for run_nml/iforcing = 3 (NWP) .AND. itopo = 1 .AND. lart=TRUE	
lrad_aero_diag	J	FALSE.		writes actual aerosol optical properties to output	

Scope	$run_nml/iforcing=2$	(ECHAM)			
Default Unit Description	Select dynamic greenhouse gases scenario (read run_nml/iforcing=2	from file)	0 : select default gas volume mixing ratios -	1990 values (CMIP5)	1 : transient CMIP5 scenario from file
Unit					
Default	0				
$\mid ext{Type}$	I				
Parameter	ighg				

Defined and used in: src/namelists/mo_radiation_nml.f90

Parameter	Type	Default	Unit	Description S	Scope
nsteps	I	666-		Number of time steps of this run. Allowed	
				range is ≥ 0 ; setting a value of 0 allows writing	
				initial output (including internal remapping)	
				without calculating time steps.	
dtime	R	0.009	SO.	time step.	
				For real case runs the maximum allowable time	
				step can be estimated as	
				$1.8 \cdot \text{ndyn_substeps} \cdot \overline{\Delta x} \text{s km}^{-1},$	
				where $\overline{\Delta x}$ is the average resolution in km and	
				ndyn_substeps is the number of dynamics	
				substeps set in nonhydrostatic_nml.	
				ndyn_substeps should not be increased beyond	
				the default value 5.	
Itestcase	L	TRUE.		Idealized testcase runs	
ldynamics	Г	TRUE.		Compute adiabatic dynamic tendencies	

Parameter	Type	Default	Unit	Description	Scope
iforcing	Ι	0		Forcing of dynamics and transport by parameterized processes. Use positive indices for the ocean. 0: no forcing 1: Held-Suarez forcing 2: ECHAM forcing 3: NWP forcing 4: local diabatic forcing with physics 5: local diabatic forcing (to be done)	
ltransport ntracer	I L	.FALSE. 0		Compute large-scale tracer transport Number of advected tracers handled by the	
lvert_nest	П	FALSE.		If set to .true. vertical nesting is switched on (i.e. variable number of vertical levels)	
num_lev	I(max_dom)	31		Number of full levels (atm.) for each domain	$lvert_nest=.TRUE.$
nshift	I(max_ dom)	0		vertical half level of parent domain which coincides with upper boundary of the current domain required for vertical refinement, which is not yet implemented	lvert_nest=.TRUE.
ltimer	IJ	.TRUE.		TRUE: Timer for monitoring the runtime of specific routines is on $(FALSE = off)$	
timers_level activate_sync_timers	гп	ㅡ ഥ		TRUE: Timer for monitoring runtime of	
msg_level	I	10		communication routines (FALSE = off) controls how much printout is written during runtime.	
msg_timestamp	L	.FALSE.		For values less than 5, only the time step is written. If .TRUE., precede output messages by time stamp.	

Parameter	Type	Default	Unit	Description	Scope
test_mode	П	0		Setting a value larger than 0 activates a dummy mode in which time stepping is changed into just doing iterations, and MPI communication is replaced by copying some value from the send buffer into the receive	iequations $= 3$
debug_check_level	Ι	0		buffer (does not work with nesting and reduced radiation grid because the send buffer may then be empty on some PEs) Setting a value larger than 0 activates debug checks.	
output	C(:)	"nml", "totint"		Main switch for enabling/disabling components of the model output. One or more choices can be set (as an array of string constants). Possible choices are:	
				• "none": switch off all output;	
				• "nml": new output mode (cf. output_nml);	
				• "totint": computation of total integrals.	
				• "maxwinds": write max. winds to separate ASCII file "maxwinds.log".	
				If the output namelist parameter is not set explicitly, the default setting "nml", "totint" is assumed.	
restart_filename	D			File name for restart/checkpoint files (containing keyword substitution patterns <gridfile>, <idom>, <rsttime>, <mtype>). default:</mtype></rsttime></idom></gridfile>	
				" <pre>"<gridfile>_restart_<mtype>_<rsttime>.nc".</rsttime></mtype></gridfile></pre>	

escription Scope	controls how profiling printout is written: TIMER_MODE_AGGREGATED=1, TIMER_MODE_DETAILED=2, TIMER_MODE_WRITE_FILES=3.	Main switch which enables the treatment of atmospheric aerosol and trace gases (The ART package of KIT is needed for this purpose)	If this flag is set to .TRUE. we give only
Unit]			
Default Unit Description	1	.FALSE.	FALSE.
Type	I	П	Г
Parameter	profiling_output	lart	check_uuid_gracefully

Defined and used in: src/namelists/mo_run_nml.f90

4.4.32 sleve_nml (relevant if nonhydrostatic_nml:ivctype=2)

Parameter	Type	Default	Unit	Description	Scope
min_lay_thckn	m R	50	ш	Layer thickness of lowermost layer; specifying	
				zero or a negative value leads to constant layer	
				thicknesses determined by top_height and nlev	
max_lay_thckn	R	25000	m	Maximum layer thickness below the height	
				given by htop_thcknlimit (NWP	
				recommendation: 400 m)	
				Use with caution! Too ambitious settings may	
				result in numerically unstable layer	
				configurations.	
htop_thcknlimit	m R	15000	m	Height below which the layer thickness does	
				not exceed max_lay_thckn	
itype_laydistr	I	1		Type of analytical function used to specify the	
				distribution of the vertical coordinate surfaces	
				1: transformed cosine, 2: third-order	
				polynomial	
top_height	$_{ m R}$	23500.0	m	Height of model top	

Parameter	Type	Default Unit		Description Scope	ope
stretch_fac	m R	1.0		Stretching factor to vary distribution of model	
				levels; values <1 increase the layer thickness	
				near the model top	
decay_scale_1	R	4000	ш	Decay scale of large-scale topography	
				component	
decay_scale_2	R	2500	m	Decay scale of small-scale topography	
				component	
decay_exp	R	1.2		Exponent of decay function	
flat_height	R	16000	m	Height above which the coordinate surfaces are	
				flat	
lread_smt	П	.FALSE.		read smoothed topography from file (TRUE)	
				or compute internally (FALSE)	

Defined and used in: src/namelists/mo_sleve_nml.f90

4.4.33 synsat_nml1

This namelist enables the RTTOV library incorporated into ICON for simulating satellite radiance and brightness temperatures. RTTOV is a radiative transfer model for nadir-viewing passive visible, infrared and microwave satellite radiometers, spectrometers and interferometers,

https://nwpsaf.eu/deliverables/rtm

for detailed information.

Parameter	Type	Default	Unit	Description	Scope
lsynsat	Γ	FALSE.		Main switch: Enables/disables computation of	
	(max-dom			synthetic satellite imagery for each model	
				domain.	
nlev_rttov	Ι	51		Number of RTTOV levels.	

¹Important note: This feature is currently active for configuration dwd+cray only.

Enabling the synsat module makes the following 32 two-dimensional output fields available:

SYNMSG_RAD_CL_IR3.9	SYNMSG_BT_CL_IR3.9	SYNMSG_RAD_CL_WV6.2	SYNMSG_BT_CL_WV6.2
SYNMSG_RAD_CL_WV7.3	SYNMSG_BT_CL_WV7.3	SYNMSG_RAD_CL_IR8.7	SYNMSG_BT_CL_IR8.7
SYNMSG_RAD_CL_IR9.7	SYNMSG_BT_CL_IR9.7	SYNMSG_RAD_CL_IR10.8	SYNMSG_BT_CL_IR10.8
SYNMSG_RAD_CL_IR12.1	SYNMSG_BT_CL_IR12.1	SYNMSG_RAD_CL_IR13.4	SYNMSG_BT_CL_IR13.4
SYNMSG_RAD_CS_IR3.9	SYNMSG_BT_CS_IR3.9	SYNMSG_RAD_CS_WV6.2	SYNMSG_BT_CS_WV6.2
SYNMSG_RAD_CS_WV7.3	SYNMSG_BT_CS_WV7.3	SYNMSG_RAD_CS_IR8.7	SYNMSG_BT_CS_IR8.7
SYNMSG_RAD_CS_IR9.7	SYNMSG_BT_CS_IR9.7	SYNMSG_RAD_CS_IR10.8	SYNMSG_BT_CS_IR10.8
SYNMSG_RAD_CS_IR12.1	SYNMSG_BT_CS_IR12.1	SYNMSG_RAD_CS_IR13.4	SYNMSG_BT_CS_IR13.4

Here, RAD denotes radiance, BT brightness temperature, CL cloudy, and CS clear sky, supplemented by the channel name.

Defined and used in: src/namelists/mo_synsat_nml.f90

4.4.34 time_nml

arameter	m Type	Default	Unit	Description	Scope
calendar	П	1		Calendar type:	
				0=Julian/Gregorian	
				1=proleptic Gregorian	
				2=30day/month, 360day/year	

Scope	This ong the to a file and a resumed, d of time can and and litting from an is longer by the east one restart ille will not a t file will not a tart cycle.	tion	no	ith step 0 ndard run or
Description	Length of restart cycle in seconds. This namelist parameter specifies how long the model runs until it saves its state to a file and stops. Later, the model run can be resumed, s.t. a simulation over a long period of time can be split into a chain of restarted model runs. Note that the frequency of writing restart files is controlled by io_nml:dt_checkpoint. Only if the value of dt_checkpoint resulting from model default or user's specification is longer than dt_restart, it will be reset (by the model) to dt_restart so that at least one restart file is generated during the restart cycle. If dt_restart is larger than but not a multiple of dt_checkpoint, restart file will not be generated at the end of the restart cycle.	Initial date and time of the simulation	End date and time of the simulation	TRUE., if time loop shall start with step 0 regardless whether we are in a standard run or
Unit	ω			
Default	86400.*30.	,2008- 09-01T 00:00:00Z'	'2008- 09-01T 01:40:00Z'	.FALSE.
Type	ಆ	O	D	Г
Parameter	dt_restart	ini_datetime_string	end_datetime_string	is_relative_time

Length of the run If "nsteps" in run_nml is positive, then nsteps*dtime is used to compute the end date and time of the run. Else the initial date and time, the end date and time, dt_restart, as well as the time step are used to compute "nsteps".

4.4.35 transport_nml (used if run_nml/ltransport=.TRUE.)

Parameter	Type	Default	Unit	Description	Scope
lvadv_tracer	Г	.TRUE.		TRUE: compute vertical tracer advection	
				FALSE: do not compute vertical tracer	
				advection	
ihadv_tracer	I(ntracer)	2		Tracer specific method to compute horizontal	
				advection:	
				0: no horiz. transport (note that the specific	
				tracer quantity q is kept constant and not	
				tracer mass ρq)	
				1: upwind (1st order)	
				2: Miura (2nd order, linear reconstr.)	
				3: Miura3 (quadr. or cubic reconstr.)	$lsq_high_ord \in [2,3]$
				4: FFSL (quadr. or cubic reconstr.)	$lsq_high_ord \in [2,3]$
				5: hybrid Miura3/FFSL (quadr. or cubic	$lsq_high_ord \in [2,3]$
				reconstr.)	
				20: miura (2nd order, lin. reconstr.) with	
				subcycling	
				22: combination of miura and miura with	
				subcycling	
				32: combination of miura and miura with	
				subcycling	
				42: combination of FFSL and miura with	
				subcycling	
				52: combination of hybrid FFSL/Miura3 with	
				subcycling	
				Subcycling means that the integration from	
				time step n to n+1 is splitted into substeps to	
				meet the stability requirements. For NWP	
				runs, substepping is generally applied above	
				$z = 22 \mathrm{km}$ (see	
				$nonhydrostatic_nml/hbot_qvsubstep).$	

Parameter	Type	Default	Unit	Description	Scope
ivadv_tracer	I(ntracer)	3		Tracer specific method to compute vertical	lvadv_tracer=TRUE
				advection:	
				0: no vert. transport (note that tracer mass ρq	
				instead of the specific tracer quantity q is kept	
				constant. This differs from the behaviour in	
				horizontal direction!)	
				1: upwind (1st order)	
				3: ppm_cfl (3^{rd} order, handles CFL > 1)	
				30: ppm (3rd order, $CFL \le 1$)	
iadv-tke	Ι	0		Type of TKE advection	inwp_turb=1
				0: no TKE advection	
				1: vertical advection only	
				2: vertical and horizontal advection	
lstrang	Γ	.FALSE.		Time splitting method	
				TRUE: second order Strang splitting	
				FALSE: first order Godunov splitting	
ctracer_list	C	r.		Two purposes:	
				- used for selecting those tracers which should	[nh/ha]_test_name=
				be initialized for idealized testcases.	'jabw','PA','DF'
				- used for tracer output names. In some	iforcing≠ inwp, iecham
				idealized cases tracers are named 'Qx', with x	
				being a 1-digit integer taken from	
				ctracer_list.	
npassive_tracer	Ι	0		number of additional passive tracers which	
				have no sources and are transparent to any	
				physical process (no effect).	
				Passive tracers are named Qpassive_ID, where	
				ID is a number between ntracer and	
				ntracer+npassive_tracer.	
				NOTE: By default, limiters are switched of for	
				passive tracers and the scheme 52 is selected	
				for horizontal advection.	

Parameter	Type	Default	Unit	Description	Scope
init_formula	C	2.2		Comma-separated list of initialization formulas	npassive_tracer; 0
				for additional passive tracers.	
itype_hlimit	I(ntracer)	4		Type of limiter for horizontal transport:	
				0: no limiter	
				3: monotonous flux limiter	
				4: positive definite flux limiter	
itype_vlimit	I(ntracer)			Type of limiter for vertical transport:	
				0: no limiter	
				1: semi-monotone slope limiter	
				2: monotonous slope limiter	
				4: positive definite flux limiter	
beta_fct	R	1.005		factor of allowed over-/undershooting in	$itype_hlimit = 3$
				monotonous limiter	
iord_backtraj	I	1		order of backward trajectory calculation:	
				1: first order	
				2: second order (iterative; currently 1 iteration	ihadv_tracer='miura'
				hardcoded; experimental!)	
igrad_c_miura	ı			Method for gradient reconstruction at cell	
				center for 2nd order miura scheme	
				1: Least-squares (linear, non-consv)	$ihadv_tracer=2$
				2: Green-Gauss	
ivcfl_max	I	ಬ		determines stability range of vertical	ivadv_tracer=3
				PPM-scheme in terms of the maximum	
				allowable CFL-number	
llsq_svd	ı	TRUE.		use QR decomposition (FALSE) or SV	
				decomposition (TRUE) for least squares design	
				matrix A	
lclip_tracer	Γ	FALSE.		Clipping of negative values	
				-	

Defined and used in: src/namelists/mo_advection_nml.f90

4.4.36 turbdiff_nml

Parameter	Type	Default	Unit	Description	Scope
imode_turb	I	1		Mode of solving the TKE equation for	
				atmosph. layers:	
				0: diagnostic equation	
				1: prognostic equation (current version)	
				2: prognostic equation (intrinsically positive	
				definite)	
imode_tran	Ι	0		Same as <i>imode_turb</i> but only for the transfer	
				layer	
icldm_turb	I	2		Mode of water cloud representation in	
				turbulence for atmosph. layers:	
				-1: ignoring cloud water completely (pure dry	
				scheme)	
				0: no clouds considered (all cloud water is	
				evaporated)	
				1: only grid scale condensation possible	
				2: also sub grid (turbulent) condensation	
				considered	
icldm_tran	Ι	2		Same as <i>icldm_turb</i> but only for the transfer	
				layer	
itype_wcld	I	2		type of water cloud diagnosis within the	icldm_turb=2 or
				turbulence scheme:	icldm_tran=2
				1: employing a scheme based on relative	
				humitidy	
				2: employing a statistical saturation adjustment	
itype_sher	I	0		Type of shear forcing used in turbulence:	
				0: only vertical shear of horizontal wind	
				1: previous plus horizontal shear correction	
				2: previous plus shear from vertical velocity	
				3: same as option 1, but (when combined with	
				ltkeshs=.TRUE.) scaling of coarse-grid	
				horizontal shear production term with $\frac{1}{\sqrt{Ri}}$	

Parameter	Type	Default	Unit	Description	Scope
ltkeshs	I	.FALSE.		Include correction term for coarse grids in	itype_sher ≥ 1
				horizontal shear production term (needed at	
				non-convection-resolving model resolutions in	
	١	ļ			,
Itkesso	コ	FALSE.		der TKE-production by sub grid SSO	$lnwp_so = 1$
ltkecon	Ļ	FALSE		wakes Consider TKE-production by sub grid	$\lim_{n\to\infty} conv = 1$
	1	2			
Itkeshs	Γ	FALSE.		Consider TKE-production by separated	
				horizontal shear eddies (inactive)	
ltmpcor	J	FALSE.		Consider thermal TKE sources in enthalpy	
				equation	
lsflend	Г	TRUE.		Use lower flux condition for vertical diffusion	
				calculation (TRUE) instead of a lower	
				concentration condition (FALSE)	
lexpcor	Γ	FALSE.		Explicit corrections of implicitly calculated	
				vertical diffusion of non-conservative scalars	
				that are involved in sub grid condensation	
				processes	
turlen	$_{ m R}$	500.0	m	Asymptotic maximal turbulent distance	
				$(\kappa * tur_len$ is the integral turbulent master	
				length scale)	
pat_len	$_{ m R}$	100.0	m	Effective length scale of thermal surface	
				patterns controlling TKE-production by sub	
				grid kata/ana-batic circulations. In case of	
				$pat_len = 0$, this production is switched off.	
c_diff	$_{ m R}$	0.2		Length scale factor for vertical diffusion of	
				TKE. In case of $cdiff = 0$, TKE is not	
				diffused vertically.	
a_stab	$_{ m R}$	0.0	1	Factor for stability correction of turbulent	
				length scale. In case of $a_stab = 0$, the	
				turbulent length scale is not reduced for stable	
				stratification.	

Parameter	Type	Default	Unit	Description	Scope
a_hshr	R	0.20	1	Length scale factor for the separated horizontal	ltkeshs=.TRUE.
				shear mode. In case of $ahshr = 0$, this shear	
alpha0	R	0.0123		Lower bound of velocity-dependent Charnock	
-	۲	0		parameter	
a.pnau_max	Υ.	0.0335	_	Upper bound of velocity-dependent Charnock parameter. Setting this parameter to 0.0335 or	
				higher values implies unconstrained velocity	
				dependence	
tkhmin	R	0.75	m^2/s	Scaling factor for minimum vertical diffusion	
				coefficient (proportional to $1/\sqrt{Ri}$) for heat	
tkmmin	22	0.75	m ² /s	Scaling factor for minimum vertical diffusion	
	2) • • •		coefficient (proportional to $1/\sqrt{Ri}$) for	
				momentum	
tkmmin_strat	${ m R}$	ರ	m^2/s	Enhanced scaling factor for minimum vertical	
				diffusion coefficient (proportional to $1/\sqrt{Ri}$)	
				for momentum, valid in the stratosphere above	
				30 km	
tkhmin_strat	\mathbf{R}	ಬ	m^2/s	Enhanced scaling factor for minimum vertical	
				diffusion coefficient (proportional to $1/\sqrt{Ri}$)	
				for heat and moisture, valid in the stratosphere	
				above 30 km	
itype_synd	I	2		Type of diagnostics of synoptic near surface	
				variables:	
				1: Considering the mean surface roughness of a	
				grid box	
				2: Considering a fictive surface roughness of a SYNOP lawn	
rlam_heat	\mathbb{R}	1.0		Scaling factor of the laminar boundary layer	
				for heat (scalars). The larger rlam heat, the larger is the laminar resistance.	

Parameter	Type	Default	Unit	Description	Scope
rat_sea	R	10.0	1	Ratio of laminar scaling factors for scalars over	
				sea and land. The larger rat_sea, the larger is	
				the laminar resistance for a sea surface	
				compared to a land surface.	
tkesmot	R	0.15	1	Time smoothing factor within [0,1] for TKE.	
				In case of $tkesmot = 0$, no smoothing is active.	
fresmot	R	0.0	1	Vertical smoothing factor within [0, 1] for TKE	
				forcing terms. In case of $fremot = 0$, no	
				smoothing is active.	
imode_frcsmot	I	1		1 = apply vertical smoothing (if frcsmot>0)	
				uniformly over the globe	
				2 = restrict vertical smoothing to the tropics	
				(reduces the moist bias in the tropics while	
				avoiding adverse effects on NWP skill scores in	
				the extratropics)	
impl_s	R	1.20	1	Implicit weight near the surface (maximal	
				value)	
implt	R	0.75		Implicit weight near top of the atmosphere	
				(minimal value)	
lconst_z0	ı	FALSE.		TRUE: horizontally homogeneous roughness	
				length z0	
const_z0	$_{ m R}$	0.001	m	value for horizontally homogeneous roughness	lconst_z0=.TRUE.
				length z0	
ldiff_qi	J	FALSE.		Turbulent diffusion of cloud ice, if .TRUE.	
itype_tran	I	2		type of surface-atmosphere transfer	
Iprfcor	ı	.FALSE.		using the profile values of the lowest main level	
				instead of the mean value of the lowest layer	
				for surface flux calculations	
Inonloc	Г	FALSE.		nonlocal calculation of vertical gradients used	
				for turbul. diff.	

Parameter Ifreeslip	Type L	Default .FALSE.	Unit	Default Unit Description FALSE. TRUE: use a free-slip lower boundary condition, i.e. neither momentum nor	Scope
	L	.FALSE.		heat/moisture fluxes (use for idealized runs only!) consideration of fluctuations of the heat capacity of air	

Defined and used in: src/namelists/mo_turbdiff_nml.f90

4.4.37 vdiff_nml

Parameter	Type	Default	Unit	Description	Scope
lsfc_mon_flux	L	.TRUE.		Switch on surface momentum flux.	lvdiff = .TRUE.
lsfc_heat_flux	П	TRUE.		Switch on surface sensible and latent heat flux.	lvdiff = .TRUE.

Defined and used in: src/namelists/mo_vdiff_nml.f90

4.5 Ocean-specific namelist parameters

4.5.1 ocean_physics_nml

Parameter	Type	Default	Unit	Unit Description	Scope
i_sea_ice	I	1		0: No sea ice, 1: Include sea ice	
				.FALSE.: compute drag only	
richardson_factor_tracer	Ι	0.5e-5	s/m		
richardson_factor_veloc	Ι	0.5e-5	m/s		
l_constant_mixing	Τ	.FALSE.			

4.5.2 sea_ice_nml (relevant if run_nml/iforcing=2 (ECHAM))

i.ice_therm I 2 i.ice_dyn I 0 i.ice_albedo I 1		Switch for thermodynamic model: 1: Zero-layer model 2: Two layer Winton (2000) model	In an ocean run
I		1: Zero-layer model 2: Two layer Winton (2000) model	T
I I		2: Two layer Winton (2000) model	1_sea_ice must be $i=1$.
I I			In an atmospheric run
I I		3: Zero-layer model with analytical forcing (for	the ice surface type
I I		diagnostics)	must be defined.
I I		4: Zero-layer model for atmosphere-only runs	
Ι		(for diagnostics)	
i_ice_albedo I 1		Switch for sea-ice dynamics:	
i_ice_albedo I 1		0: No dynamics	
i_ice_albedo I I 1		1: FEM dynamics (from AWI)	
		Switch for albedo model. Only one is	
		implemented so far.	
i.Qio_type		Switch for ice-ocean heat-flux calculation	Defaults to 1 when
		method:	i.ice_dyn=0 and 2
		1: Proportional to ocean cell thickness (like	otherwise.
		MPI-OM)	
		2: Proportional to speed difference between ice	
		and ocean	
		Number of ice classes (must be one for now)	
Я.	 m	Hibler's h_0 parameter for new-ice growth.	
	m	Minimum sea-ice thickness allowed.	
	days	Number of days it takes the wind to reach	
		correct strength. Only used at the start of an	
		OMIP/NCEP simulation (not after restart).	

4.6 Namelist parameters for testcases (NAMELIST_ICON)

The ICON model code includes several experiments, so-called test cases, for the shallow water model as well as the 3-dimensional atmosphere. Depending on the specified experiment, initial conditions and boundary conditions are computed internally.

4.6.1 ha_testcase_nml (Scope: Itestcase=.TRUE. and iequations=[0,1,2] in run_nml)

Parameter	Type	Default	Unit	Description	Scope
ctest_name	C	'JWw'		Name of test case:	
				'SW_GW': gravity wave	lshallow_water=.TRUE.
				'USBR': unsteady solid body rotation	lshallow_water=.TRUE.
				'Will_2': Williamson test 2	lshallow_water=.TRUE.
				'Will_3': Williamson test 3	lshallow_water=.TRUE.
				'Will_5': Williamson test 5	lshallow_water=.TRUE.
				'Will_6': Williamson test 6	lshallow_water=.TRUE.
				'GW': gravity wave ($nlev=20 only!$)	lshallow-water=.FALSE.
				'LDF': local diabatic forcing test without	lshallow_water=.FALSE.
				physics	and iforcing=4
				'LDF-Moist': local diabatic forcing test with	lshallow_water=.FALSE.,
				physics initalised with zonal wind field	and iforcing=5
				'HS': Held-Suarez test	lshallow_water=.FALSE.
				'JWs': Jablonowski-Will. steady state	lshallow_water=.FALSE.
				'JWw': Jablonowski-Will. wave test	lshallow_water=.FALSE.
				'JWw-Moist': Jablonowski-Will. wave test	lshallow_water=.FALSE.
				including moisture	
				'APE': aqua planet experiment	lshallow_water=.FALSE.
				'MRW': mountain induced Rossby wave	lshallow_water=.FALSE.
				'MRW2': modified mountain induced Rossby	lshallow_water=.FALSE.
				Wave	
				'PA': pure advection	lshallow_water=.FALSE.
				'SV': stationary vortex	lshallow_water=.FALSE.,
					ntracer = 2
				'DF1': deformational flow test 1	
				'DF2': deformational flow test 2	
				'DF3': deformational flow test 3	
				'DF4': deformational flow test 4	
				'RH': Rossby-Haurwitz wave test	lshallow_water=.FALSE.

Parameter	Type	Default	Unit	Description	Scone
rotate axis deg	B	0.0	deg	Farth's rotation axis pitch angle	ctest name= 'Will 2'.
0	, I)	0		Will_3', 'JWs', 'JWw',
					'PA', 'DF1234'
gw_brunt_vais	R	0.01	1/s	Brunt Vaisala frequency	ctest_name= 'GW'
gw_u0	R	0.0	$\mathrm{m/s}$	zonal wind parameter	ctest_name= 'GW'
gw_lon_deg	Я	180.0	\deg	longitude of initial perturbation	ctest_name= 'GW'
gw-lat-deg	R	0.0	deg	latitude of initial perturbation	ctest_name= 'GW'
jw_uptb	R	1.0	s/m	amplitude of the wave pertubation	ctest_name= 'JWw'
			(3)		
mountctr_lon_deg	R	0.06	\deg	longitude of mountain peak	ctest_name= 'MRW(2)'
mountctr_lat_deg	R	30.0	\deg	latitude of mountain peak	ctest_name= 'MRW(2)'
mountctr_height	Я	2000.0	m	mountain height	ctest_name= 'MRW(2)'
mountctr_half_width	Я	1500000.0	m	mountain half width	ctest_name= 'MRW(2)'
mount_u0	R	20.0	$\mathrm{m/s}$	wind speed for MRW cases	ctest_name= 'MRW(2)'
rh_wavenum	I	4		wave number	ctest_name= 'RH'
rh_init_shift_deg	R	0.0	\deg	pattern shift	ctest_name= 'RH'
ihs_init_type	I	1		Choice of initial condition for the Held-Suarez	ctest_name= 'HS'
				test. 1: the zonal state defined in the JWs test	
				case; other integers: isothermal state	
				(T=300 K, ps=1000 hPa, u=v=0.)	
lhs_vn_ptb	L	TRUE.		Add random noise to the initial wind field in	ctest_name= 'HS'
				the Held-Suarez test.	
hs_vn_ptb_scale	R	÷	$\mathrm{m/s}$	Magnitude of the random noise added to the	ctest_name= 'HS'
				initial wind field in the Held-Suarez test.	
lrh_linear_pres	L	FALSE.		Initialize the relative humidity using a linear	ctest_name=
				function of pressure.	'JWw-Moist', 'APE',
					'LDF-Moist'
$rh_at_1000hpa$	R	0.75		relative humidity	ctest_name=
				0,1	'JWw-Moist', 'APE',
				at 1000 hPa	'LDF-Moist'
linit_tracer_fv	L	TRUE.		Finite volume initialization for tracer fields	ctest_name='PA'

Parameter	Type	Default	Unit	Description	Scope
ape_sst_case	C	'sst1'		SST distribution selection	ctest_name='APE'
				'sst1': Control experiment	
				'sst2': Peaked experiment	
				'sst3': Flat experiment	
				'sst4': Control-5N experiment	
				'sst_qobs': Qobs SST distribution exp	
				'sst_ice': Control SST distribution with -1.8 C	
				above 64 N/S .	
ildf_init_type	I	0		Choice of initial condition for the Local	$ctest_name = 'LDF'$
				diabatic forcing test. 1: the zonal state defined	
				in the JWs test case; other: isothermal state	
				(T=300 K, ps=1000 hPa, u=v=0.)	
ldf_symm	Γ	TRUE.		Shape of local diabatic forcing:	ctest_name=
				.TRUE.: local diabatic forcing symmetric	'LDF', 'LDF-Moist'
				about the equator (at 0 N)	
				.FALSE.: local diabatic forcing asym. about	
				the equator (at 30 N)	

Defined and used in: src/testcases/mo_ha_testcases.f90

4.6.2 nh_testcase_nml (Scope: Itestcase=.TRUE. and iequations=3 in run_nml)

Parameter	Type	Default	Unit	Default Unit Description	Scope
nh_test_name	C	'jabw'		testcase selection	
				'zero': no orography	
				'bell': bell shaped mountain at 0E,0N	
				'schaer': hilly mountain at 0E,0N	
				'jabw': Initializes the full Jablonowski	
				Williamson test case.	
				'jabw_s': Initializes the Jablonowski	
				Williamson steady state test case.	

_																																
Scope																									$llimited_area = .TRUE.$							
Description	'jabw_m': Initializes the Jablonowski	Williamson test case with a mountain instead	of the wind perturbation (specify	mount_height).	'mrw_nh': Initializes the full	Mountain-induced Rossby wave test case.	'mrw2_nh': Initializes the modified	mountain-induced Rossby wave test case.	'mwbr_const': Initializes the mountain wave	with two layers test case. The lower layer is	isothermal and the upper layer has constant	brunt vaisala frequency. The interface has	constant pressure.	' PA ': Initializes the pure advection test case.	'HS_nh': Initializes the Held-Suarez test case.	At the moment with an isothermal atmosphere	at rest (T=300K, ps=1000hPa, u=v=0,	topography=0.0).	'HS_jw': Initializes the Held-Suarez test case	with Jablonowski Williamson initial conditions	and zero topography.	'APE_nwp, APE_echam, APE_nh,	APEc_nh , ': Initializes the APE experiments.	With the jabw test case, including moisture.	'wk82': Initializes the Weisman Klemp test	case	'g_lim_area': Initializes a series of general	limited area test cases: itype_atmos_ana	determines the atmospheric profile,	itype_anaprof_uv determines the wind profile	and itype_topo_ana determines the topography	'dcmip_bw_ll': initializes (moist) baroclinic instability/wave (DCMIP2016)
Unit																																
Default																																
Type																																
Parameter																																

Parameter	Type	Default	Unit	Description	Scope
	4			'dcmip_pa_12': Initializes Hadley-like meridional circulation pure advection test case.	4
				'dcmip_rest_200': atmosphere at rest test (Schaer-type mountain)	lcoriolis = .FALSE.
	-			'dcmip_mw_2x': nonhydrostatic mountain waves triggered by Schaer tyne mountain	lcoriolis = .FALSE.
				'dcmip_gw_31': nonhydrostatic gravity waves	
				triggered by a localized perturbation (nonlinear)	
				'dcmip_gw_32': nonhydrostatic gravity waves	llimited_area =.TRUE.
				'demip_tc_51': tropical cyclone test case with	lcoriolis = .TRUE.
				'simple physics' parameterizations (not yet	
				implemented)	
				$'$ dcmip_tc_52 $'$: tropical cyclone test case with	lcoriolis = .TRUE.
				with full physics in Aqua-planet mode	
				'CBL': convective boundary layer simulations	is_plane_torus=
				for LES package on torus (doubly periodic) grid	TRUE.
is_toy_chem	П	.FALSE.		Terminator toy chemistry activated when	
				TRUE.	
$\operatorname{dcmip_bw\%}$				DCMIP2016 baroclinic wave test	'dcmip_bw_11'
deeb	I	0		deep atmosphere	
				(1 = yes or 0 = no)	
moist	I	0		include moisture, i.e. $qv \neq 0$	
				(1 = yes or 0 = no)	
pertt	I	0		type of initial perturbation	
				(0 = exponential, 1 = stream function)	
${ m toy_chem\%}$				terminator toy chemistry	is_toy_chem=.TRUE.
dt-chem	m R	300	\mathbf{S}	chemistry tendency update interval	
$dt_{-}cpl$	Я	300	ß	chemistry-transport coupling interval	
id_cl	I	1		Tracer container slice index for species CL	
id_cl2	I	2		Tracer container slice index for species CL2	

Parameter	Type	Default	Unit	Description	Scope
du-wį	R	1.0	s/w	amplitude of the u-perturbation in jabw test	nh_test_name='jabw'
				case	
u0_mrw	$_{ m R}$	20.0	s/m	wind speed for $mrw(2)$ and $mwbr_const$ cases	nh_test_name=
					$\operatorname{mrw}(2)$ _nh' and
					'mwbr_const'
mount_height_mrw	\mathbf{R}	2000.0	m	maximum mount height in $mrw(2)$ and	nh_test_name=
				mwbr_const	'mrw(2)_nh' and
					'mwbr_const'
mount_half_width	m R	1500000.0	m	half width of mountain in $mrw(2)$, $mwbr_{-const}$	nh_test_name=
				and bell	$\operatorname{mrw}(2)$ -nh',
					'mwbr_const' and 'bell'
mount_lonctr_mrw_deg	$_{ m R}$	90.	deg	lon of mountain center in $mrw(2)$ and	nh_test_name=
				mwbr_const	$\operatorname{mrw}(2)$ _nh' and
					'mwbr_const'
mount_latctr_mrw_deg	\mathbf{R}	30.	deg	lat of mountain center in $mrw(2)$ and	nh_test_name=
				mwbr_const	'mrw(2)_nh' and
					'mwbr_const'
temp_i_mwbr_const	\mathbf{R}	288.0	X	temp at isothermal lower layer for mwbr_const	nh_test_name=
				case	'mwbr_const'
p_int_mwbr_const	\mathbf{R}	70000.	Pa	pres at the interface of the two layers for	nh_test_name=
				mwbr_const case	'mwbr_const'
bruntvais_u_mwbr_const	\mathbf{R}	0.025	1/s	constant brunt vaissala frequency at upper	nh_test_name=
				layer for mwbr_const case	'mwbr_const'
mount_height	\mathbf{R}	100.0	m	peak height of mountain	nh_test_name= 'bell'
layer_thickness	$_{ m R}$	-999.0	m	thickness of vertical layers	If layer_thickness < 0 ,
					the vertical level
					distribution is read in
					from externally given
					HYB_PARAMS_XX.
n_flat_level	П	2		level number for which the layer is still flat and	layer_thickness > 0
				not terrain-following	
np-n0	H .	0.0	s/m	initial constant zonal wind speed	nh_test_name = 'bell'
$\mid \mathrm{nh}_{-}\mathrm{t0} \mid$	\mathbb{R}	300.0	X	initial temperature at lowest level	$ \text{nh_test_name} = \text{'bell'} $

Parameter	Type	Default	Unit	Description	Scope
nh_brunt_vais	R	0.01	1/s	initial Brunt-Vaisala frequency	nh_test_name = 'bell'
torus_domain_length	R	100000.0	m	length of slice domain	$nh_test_name = 'bell',$
					lplane=.TRUE.
rotate_axis_deg	В	0.0	deg	Earth's rotation axis pitch angle	nh_test_name= 'PA'
lhs_nh_vn_ptb	Г	TRUE.		Add random noise to the initial wind field in	nh_test_name= 'HS_nh'
				the Held-Suarez test.	
lhs_fric_heat	T	FALSE.		add frictional heating from Rayleigh friction in	nh_test_name= 'HS_nh'
				the Held-Suarez test.	
hs_nh_vn_ptb_scale	R	1.	s/m	Magnitude of the random noise added to the	nh_test_name= 'HS_nh'
				initial wind field in the Held-Suarez test.	
rh_at_1000hpa	R	0.7	1	relative humidity at 1000 hPa	nh_test_name= 'jabw',
					nh_test_name= 'mrw'
qv_max	R	20.e-3	kg/kg	specific humidity in the tropics	nh_test_name= 'jabw',
					nh_test_name= 'mrw'
${ m ape_sst_case}$	C	'sst1'		SST distribution selection	$nh_test_name = APE_nwp',$
				'sst1': Control experiment	'APE_echam'
				'sst2': Peaked experiment	
				'sst3': Flat experiment	
				'sst4': Control-5N experiment	
				'sst_qobs': Qobs SST distribution exp.	
				'sst_const': constant SST	
ape_sst_val	R	29.0	$_{ m degC}$	aqua planet SST for ape_sst_case='sst_const'	nh_test_name=
					$^{\prime}\mathrm{APE_nwp'},$
					'APE_echam'
linit_tracer_fv	Γ	.TRUE.		Finite volume initialization for tracer fields	pure advection tests,
					only
lcoupled_rho	П	.FALSE.		Integrate density equation 'offline'	pure advection tests,
					only
qv_max_wk	Я	0.014	Kg/kg	maximum specific humidity near the surface, range 0.012 - 0.016 used to vary the buoyancy	nh_test_name='wk82'
	_	_			_

Parameter	Type	Default	Unit	Description	Scope
u_infty_wk	R	20.	s/m	zonal wind at infinity height	nh_test_name='wk82'
				range 0 45.	
				used to vary the wind shear	
bub_amp	R	2.	X	maximum amplitud of the thermal	nh_test_name='wk82'
				perturbation	
bubctr_lat	R	0.	\deg	latitude of the center of the thermal	nh_test_name='wk82'
				perturbation	
bubctr_lon	$_{ m R}$.06	\deg	longitude of the center of the thermal	nh_test_name='wk82'
				perturbation	
$bubctr_z$	$_{ m R}$	1400.	m	height of the center of the thermal perturbation	nh_test_name='wk82'
bub_hor_width	$_{ m R}$	10000.	m	horizontal radius of the thermal perturbation	nh_test_name='wk82'
$\mathrm{bub_ver_width}$	$_{ m R}$	1400.	m	vertical radius of the thermal perturbation	nh_test_name='wk82'
itype_atmo_ana	I	1		kind of atmospheric profile:	nh_test_name=
				1 piecewise N constant layers	'g_lim_area'
				2 piecewise polytropic layers	
itype_anaprof_uv	I	1		kind of wind profile:	nh_test_name=
				1 piecewise linear wind layers	'g_lim_area'
				2 constant zonal wind	
				3 constant meridional wind	
itype_topo_ana	I	1		kind of orography:	nh_test_name=
				1 schaer test case mountain	'g_lim_area'
				2 gaussian_2d mountain	
				3 gaussian_3d mountain	
				any other no orography	
nlayers_nconst	I	П		Number of the desired layers with a constant	nh_test_name=
				Brunt-Vaisala-frequency	'g_lim_area' and
					itype_atmo_ana=1
p_base_nconst	R	100000.	Pa	pressure at the base of the first N constant	nh_test_name=
				layer	'g_lim_area' and
					itype_atmo_ana=1
theta0_base_nconst	R	288.	K	potential temperature at the base of the first N	nh_test_name=
				constant layer	'g_lim_area' and
					itype_atmo_ana=1

Parameter	Type	Default	Unit	Description	Scope
h-nconst	R(nlayers	0.,	m	height of the base of each of the N constant	nh_test_name=
	-nconst)	1500.,		layers	'g_lim_area' and
		12000.			itype_atmo_ana=1
N_nconst	R(nlayers	0.01	1/s	Brunt-Vaisala-frequency at each of the N	nh_test_name=
	-nconst)			constant layers	'g_lim_area' and
					itype_atmo_ana=1
rh_nconst	R(nlayers	0.5	%	relative humidity at the base of each N	nh_test_name=
	-nconst)			constant layers	'g_lim_area' and
					itype_atmo_ana=1
rhgr_nconst	R(nlayers	0.	%	relative humidity gradient at each of the N	nh_test_name=
	-nconst)			constant layers	'g_lim_area' and
					itype_atmo_ana=1
nlayers_poly	Ι	2		Number of the desired layers with constant	nh_test_name=
				gradient temperature	'g_lim_area' and
					itype_atmo_ana=2
p-base-poly	$_{ m R}$	100000.	Pa	pressure at the base of the first polytropic layer	nh_test_name=
					'g_lim_area' and
					itype_atmo_ana=2
h_poly	R(nlayers	0.,	m	height of the base of each of the polytropic	nh_test_name=
	_poly)	12000.		layers	'g_lim_area' and
					itype_atmo_ana=2
t_{-} poly	R(nlayers	288.,	K	temperature at the base of each of the	nh_test_name=
	_poly)	213.		polytropic layers	'g_lim_area' and
					itype_atmo_ana=2
rh-poly	R(nlayers	0.8, 0.2	%	relative humidity at the base of each of the	nh_test_name=
	-poly)			polytropic layers	'g_lim_area' and
					itype_atmo_ana=2
rhgr-poly	R(nlayers	5.e-5, 0.	%	relative humidity gradient at each of the	nh_test_name=
	-poly)			polytropic layers	'g_lim_area' and
					itype_atmo_ana=2
nlayers_linwind	I	2		Number of the desired layers with constant U	nh_test_name=
				gradient	'g_lim_area' and
					itype_anaprof_uv=1

Parameter	Type	Default	Unit	Description	Scope
h_linwind	R(nlayers	0.,2500.	m	height of the base of each of the linear wind	nh_test_name=
	_linwind)			layers	'g_lim_area' and
		_			itype_anaprof_uv=1
u_linwind	R(nlayers	5, 10.	s/m	zonal wind at the base of each of the linear	nh_test_name=
	_linwind)	_		wind layers	'g_lim_area' and
		_			itype_anaprof_uv=1
ugr_linwind	R(nlayers	0., 0.	1/s	zonal wind gradient at each of the linear wind	nh_test_name=
	_linwind)	_		layers	'g_lim_area' and
		_			itype_anaprof_uv=1
vel_const	$_{ m R}$	20.	s/m	constant zonal/meridional wind	nh_test_name=
		_		$(itype_anaprof_uv=2,3)$	'g_lim_area' and
		_			itype_anaprof_uv=2,3
mount_lonc_deg	$_{ m R}$	90.	deg	longitud of the center of the mountain	nh_test_name=
		_			'g_lim_area'
mount_latc_deg	R	0.	deg	latitud of the center of the mountain	nh_test_name=
		_			'g_lim_area'
schaer_h0	$_{ m R}$	250.	ш	h0 parameter for the schaer mountain	nh_test_name=
		_			'g_lim_area' and
		_			itype_topo_ana=1
schaer_a	$_{ m R}$	5000.	m	-a- parameter for the schaer mountain,	nh_test_name=
		_		also half width in the north and south side of	'g_lim_area' and
		_		the finite ridge to round the sharp edges	itype_topo_ana=1,2
schaer_lambda	$_{ m R}$	4000.	m	lambda parameter for the schaer mountain	nh_test_name=
		_			'g_lim_area' and
		_			itype_topo_ana=1
lshear_dcmip	ı	FALSE		run demip_mw_2x with/without vertical wind	nh_test_name=
		_		shear	'dcmip_mw_2x'
				FALSE: dcmip_mw_21: non-sheared	
,	ı			TRUE : dcmip_mw_22: sheared	,
haltwidth_2d	H	10000.	ш	half lenght of the finite ridge in the north-south	nh_test_name=
		_		direction	'g_lim_area' and
		_			itype_topo_ana=1,2

Parameter	Type	Default	Unit	Description	Scope
m_height	R	1000.	m	height of the mountain	nh_test_name=
					'g_lim_area' and
					$itype_topo_ana=2,3$
m_width_x	${f R}$	5000.	m	half width of the gaussian mountain in the	$nh_test_name =$
				east-west direction	'g_lim_area' and
				half width in the north-south direction in the	$itype_topo_ana=2,3$
				rounding of the finite ridge (gaussian_2d)	
m_width_y	m R	5000.	m	half width of the gaussian mountain in the	$nh_test_name =$
				north-south direction	'g_lim_area' and
					$itype_topo_ana=2,3$
gw_u0	m R	0.	s/m	maximum amplitude of the zonal wind	$nh_test_name =$
					'dcmip-gw- $3X'$
gw_clat	${ m R}$	90.	deg	Lat of perturbation center	$nh_test_name =$
					'dcmip-gw- $3X'$
gw_delta_temp	\mathbb{R}	0.01	K	maximum temperature perturbation	$nh_test_name =$
					$'$ dcmip_gw_32 $'$
ucbl(2)	R	0:0	s/m	to prescribe initial zonal velocity profile for	$nh_test_name = CBL$
			and	convective boundary layer simulations where	
			1/s	$u_{-}cbl(1)$ sets the constant and $u_{-}cbl(2)$ sets the	
				vertical gradient	
vcbl(2)	m R	0:0	$\mathrm{s/m}$	to prescribe initial meridional velocity profile	$nh_test_name=CBL$
			and	for convective boundary layer simulations	
			1/s	where $v_cbl(1)$ sets the constant and $v_cbl(2)$	
				sets the vertical gradient	
$ hinspace{th_cbl(2)}$	R	290:0:006	K	to prescribe initial potential temperature	$nh_test_name = CBL$
			and	profile for convective boundary layer	
			m K/m	simulations where $thcbl(1)$ sets the constant	
				and $thcbl(2)$ sets the gradient	

Defined and used in: src/testcases/mo_nh_testcases.f90

4.7. EXTERNAL DATA 150

4.7 External data

4.7.1 extpar_nml (Scope: itopo=1 in run_nml)

Parameter	Type	Default	Unit	Description	Scope
itopo	I	0		0: analytical topography/ext. data	
				1: topography/ext. data read from file	
$n_iter_smooth_topo$	I(n-dom)	0		iterations of topography smoother	itopo = 1
fac_smooth_topo	$_{ m R}$	0.015625		pre-factor of topography smoother	$n_iter_smooth_topo > 0$
hgtdiff_max_smooth_topo	R	0.	m	RMS height difference to neighbor grid points	$n_{iter_smooth_topo} > 0$
				at which the smoothing pre-factor	
				fac_smooth_topo reaches its maximum value	
				(linear proportionality for weaker slopes)	
heightdiff_threshold	$R(n_{-}dom)$	3000.	m	height difference between neighboring grid	
				points above which additional local nabla2	
				diffusion is applied	
lrevert_sea_height	П	.FALSE.		If .TRUE., sea point heights will be reverted to	n_iter_smooth_topo ; 0
				original (raw data) heights after topography	
				smoothing was applied.	
l_emiss	ı	TRUE.		read and use external surface emissivity map	itopo = 1
extpar_filename	C			Filename of external parameter input file,	
				default: " <path>extpar_<gridfile>". May</gridfile></path>	
				contain the keyword <pre><pre>contain the keyword <pre><pre>contain</pre></pre></pre></pre>	
				substituted by model_base_dir.	
extpar_varnames_map_	C	, ,		Filename of external parameter dictionary,	
file				This is a text file with two columns separated	
				by whitespace, where left column: NetCDF	
				name, right column: GRIB2 short name. It is	
				required, if external parameter are read from a	
				file in GRIB2 format.	

Defined and used in: src/namelists/mo_extpar_nml.f90

4.8 External packages

4.9 Information on vertical level distribution

generate the terrain following coordinates. The hybrid level specification is stored in <icon home>/hyb_params/HYB_PARAMS_<nlev>. The If no vertical sleve coordinate is chosen (ivctype /=2), the hydrostatic and nonhydrostatic models need hybrid vertical level information to hydrostatic model assumes to get pressure based coordinates, the nonhydrostatic model expects height based coordinates. For further information see <icon home>/hyb_params/README.

Discussion

Document last edited by addyourname on insert date Document last edited by S Gruber on 08-01-2014.