ICON Namelist Overview

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1 ICON Namelists

1.1 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 1: Namelist files

Namelist file	Purpose	Made by script	Used by program
NAMELIST_GRAPH	Generate graphs	create_global_grids.run	grid_command
NAMELIST_GRID	Generate grids	create_global_grids.run	grid_command
NAMELIST_GRIDREF	Gen. nested domains	create_global_grids.run	grid_command
NAMELIST OCEAN GRID	Gen. ocean grid	create ocean grid.run	grid command
NAMELIST TORUS GRID	Gen. torus grid	create torus grid.run	grid command
NAMELIST ICON	Run ICON models	exp. <name>.run</name>	control model

1.2 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
- ullet 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.
- \bullet 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.

2 Namelist parameters for grid generation

2.1 Namelist parameters defining the atmosphere grid

${\bf 2.1.1} \quad {\bf graph_ini} \ ({\bf NAMELIST_GRAPH})$

Parameter	Type	Default	Unit	Description	Scope
nroot	I	2		root subdivision of initial edges	
grid_levels	I	4		number of edge bisections following the root	
				subdivision	
lplane	L	.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

2.1.2 grid_ini (NAMELIST_GRID)

Parameter	Type	Default	Unit	Description	Scope
nroot	I	2		root subdivision of initial edges	
grid_levels	I	4		number of edge bisections following the root	
				subdivision	
lplane	L	.FALSE.		switch for generating planar grid. The root level	
				consists of 8 triangles.	

Defined and used in: $src/grid_generator/mo_io_grid.f90$

${\bf 2.1.3 \quad grid_options} \; ({\bf NAMELIST_GRID})$

Parameter	Type	Default	Unit	Description	Scope
x_rot_angle	R	0.0	deg	Rotation of the icosahedron about the x-axis	
				(connecting the origin and $[0^{\circ}E, 0^{\circ}N]$)	

Parameter	Type	Default	Unit	Description	Scope
y_rot_angle	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	
				4: spring dynamics	
l_c_grid	L	.FALSE.		C-grid constraint on last level	
maxlev_optim	I	100		Maximum grid level where the optimization is	$i_type_optimize = 1 \text{ or } 4$
				applied	
beta_spring	R	0.90		tuning factor for target grid length	$i_type_optimize = 4$

Defined and used in: src/grid generator/mo io grid.f90

2.1.4 plane options (NAMELIST GRID)

Parameter	Type	Default	Unit	Description	Scope
tria arc km	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 <= x <= 1, and $-\sqrt{3}/2 <= y <= \sqrt{3}/2$. Currently the planar option can only be used as an f-plane. Defined and used in: $src/grid_generator/mo_io_grid.f90$

2.1.5 gridref_ini (NAMELIST_GRIDREF)

Parameter Type Default U	it Description	Scope	
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Parameter	Type	Default	Unit	Description	Scope
grid_root	Ι	2		root subdivision of initial edges	
start_lev	I	4		number of edge bisections following the root	
				subdivision	
n_dom	I	2		number of logical model domains, including the	
				global one	
n_phys_dom	I	n_dom		number of physical model domains, may be larger	
				than n_dom (in this case, domain merging is	
				applied)	
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 first	
	dom-1)			nested domain; needs to be specified only in case of	
				domain merging, i.e. n_dom < n_phys_dom)	
l_plot	L	.FALSE.		produces GMT plots showing the locations of the	
				nested domains	
l_circ	L	.TRUE.		Create circular (.T.) or rectangular (.F.) refined	
				domains	
l_rotate	L	.FALSE.		Rotates center point into the equator in case of	lcirc=.FALSE.
				$l_circ = .FALSE.$	
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)	
bdy_indexing_deptl	ı I	\max_{rlcell}		Number of cell rows along the lateral boundary of a	
		(=8)		model domain for which the refin_ctrl fields	
				contain the distance from the lateral boundary;	
				needs to be enlarged when lateral boundary	
				nudging is required for one-way nesting	

Parameter	Type	Default	Unit	Description	Scope
radius	R(n_dom-	30.	deg	radius of nested domain (first entry refers to first	lcirc=.TRUE.
	1)			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 refers	lcirc=.FALSE.
	1)			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 entry	lcirc=.FALSE.
	1)			refers to first nested domain; needs to be specified	
				for each nested domain separately)	
center_lon	R(n_dom-	90.	deg	center longitude of refined domain (first entry refers	
	1)			to first nested domain; needs to be specified for	
				each nested domain separately)	
center_lat	R(n_dom-	30.	deg	center latitude of refined domain (first entry refers	
	1)			to first nested domain; needs to be specified for	
				each nested domain separately)	

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

2.2 Namelist parameters defining the local grid generation

The ocean grids are created by the script run/create_ocen_grid.run

2.2.1 grid_geometry_conditions

Parameter	Type	Default	Unit	Description	Scope
no_of_conditions	I	0		Number of geometric conditions	
patch_shape	I(no_of_	0		1=rectangle; 2=circle	
	condi-				
	tions)				
patch_center_x	R(no_of	0.0	degrees	longitude of patch center	
	_ condi-				
	tions)				

Parameter	Type	Default	Unit	Description	Scope
patch_center_y	R(no_of	0.0	degrees	latitude of patch center	
	_ condi-				
	tions)				
rectangle_xradious	R(no_of_	0.0	degrees	half meridional extension of a rectangular patch	$patch_shape=1$
	condi-				
	tions)				
rectangle_yradious	R(no_of_	0.0	degrees	half zonal extension of a rectangular patch	$patch_shape=1$
	condi-				
	tions)				
circle_radious	R(no_of_	0.0	degrees	radius of a circular patch	$patch_shape=2$
	condi-				
	tions)				

Defined in mo_grid_conditions.f90

${\bf 2.2.2}\quad {\bf local_grid_optimization}$

Parameter	Type	Default	Unit	Description	Scope
use_optimization	L	.FALSE.		Apply, or not, optimization	
use_edge_springs	L	.FALSE.		Use spring dynamics	
prime_ref_length	R	1.0		Spring length coefficient	
_coeff					
use_adaptive_	L	.FALSE.		Use adaptive spring length	
spring_length					
use_local_reference	L	.FALSE.		Use locally adaptive spring length	
_length					
local_reference_	R	0.0		Coefficient of local vs global spring length	
length_coeff					
use_isotropy_force	L	.FALSE.		Use isotropy force, tends to create symmetric	
				triangles	
isotropy_rotation	R	0.0		Coefficient of the rotational isotropy force	
_coeff					

Parameter	Type	Default	Unit	Description	Scope
isotropy_stretch	R	0.0		Coefficient of the stretch isotropy force	
_coeff					
optimize_vertex	I	1		For patches the min depth of the vertices that will	
_depth				be optimized. The boundary vertices have depth 0,	
				the next level 1, etc.	

Defined in mo_local_grid_optimization.f90

${\bf 2.2.3}\quad {\bf create_ocean_grid}$

Parameter	Type	Default	Unit	Description	Scope
only_get_sea_	L	.false.		.true.:returns the whole grid with a sea-land mask;	
$land_mask$.false.:returns only the ocean grid	
smooth_ocean_	L	.true.		.true.:smooths the ocean boundaries so no triabgle	
boundary				has two boundary edges; .false.:no smoothing	
input_file	С			name of the input grid file	
elevation_file	С			name of the file containing cell elevation values for	no_of_conditions=0
				the input_file	
elevation_field	С			name of the field containing the cell elevation values	$no_of_conditions=0$
min_sea_depth	R	0.0	m	if cell elevation < min_sea_depth then the cell is	
			(nega-	consider sea	
			tive)		
set_sea_depth	R	0.0	m	if not 0, then sea cells are of set_sea_depth	
			(nega-	elevation	
			tive)		
set_min_sea_depth	R	0.0	m	if not 0, then sea cells have a maximum of	
			(nega-	set_min_sea_depth elevation	
			tive)		
edge_elev_	I	2		compute edge elevation from cells using: linear	
$interp_method$				interpolation=1; min value = 2	
output_refined_	С			name of the output refined ocean grid file	
ocean_file					

Defined in mo_create_ocean_grid.f90

2.2.4 torus grid parameters

Parameter	Type	Default	Unit	Description	Scope
y_no_of_rows	I		4	number of triangle rows of the torus grid, >=2	
x_no_of_columns	I		8	number of triangle columns of the torus grid, >=2	
edge_length	R	m	1000.0	the triangle edge length	
x_center	R	m	0.0	the x coordinate of the torus center	
y_center	R	m	0.0	the y coordinate of the torus center	
out_file_name	C			the torus grid file name	
$unfolded_torus_$	C			the unfolded torus grid file name (for plotting)	
file_name					
ascii_filename	С			the unfolded torus grid ascci file name (for plotting)	

Defined in mo_create_torus_grid.f90. See the run script run/create_torus_grid.run.

3 Namelist parameters defining the ICON 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.

3.1 master nml

Parameter	Type	Default	Unit	Description	Scope
l_restart	L	.FALSE.		If .TRUE.: Current experiment is started from a	
				restart.	
model_base_dir	С	, ,		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 will</path>	
				be substituted.	

3.2 master_model_nml (repeated for each model)

Parameter	Type	Default	Unit	Description	Scope
model_name	С			Character string for naming this component.	
model_namelist_	С			File name containing the model namelists.	
filename					
model_type	I	0		Identifies which component to run. atmosphere=1,	
				ocean=2, radiation=3, dummy_model=99	
model_min_rank	I	0		Start MPI rank for this model.	
model_max_rank	I	-1		End MPI rank for this model.	
model_inc_rank	I	0		Stride of MPI ranks.	
model_restart_info	С	restart.info		Name (including full path) of the restart info file for	
_filename				this model	

$3.3 \quad time_nml$

Parameter	Type	Default	Unit	Description	Scope
$dt_restart$	R	86400.*30.	s	Length of restart cycle in seconds. Note that the	
				frequency of writing restart files is controlled by	
				io_nml:dt_checkpoint. 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.	
calendar	Ι	1		Calendar type:	
				0=Julian/Gregorian	
				1=proleptic Gregorian	
				$2=30 ext{day/month,} 360 ext{day/year}$	
ini_datatime_string	С	'2008-09-		Initial date and time of the simulation	
		01T00:00:0	DΖ'		

Parameter	Type	Default	Unit	Description	Scope
end_datatime_string	g C	2008-09-		End date and time of the simulation	
		01T01:40:0	0Ζ'		
				Length of the run	
				If "nsteps" in run_nml (see below) 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".	

3.4 parallel_nml

Parameter	Type	Default	Unit	Description	Scope
nproma	I	1		chunk length	
n_ghost_rows	I	1		number of halo cell rows	
division_method	I	1		method of domain decomposition	
				0: read in from file	
				1: use built-in geometric subdivision	
				2: use METIS	
division_file_name	C			Name of division file	$\operatorname{division_method} = 0$
ldiv_phys_dom	L	.TRUE.		.TRUE.: split into physical domains before	$\operatorname{division_method} = 1$
				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	L	.FALSE.		.TRUE. means verification run for MPI	
				parallelization (PE 0 processes full domain)	
l_test_openmp	L	.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	
				paralllelization	

Parameter	Type	Default	Unit	Description	Scope
l_log_checks	L	.FALSE.		if .TRUE. messages are generated during each	
				synchonization step (use for debugging only)	
l_fast_sum	L	.FALSE.		if .TRUE., use fast (not	
				processor-configuration-invariant) global summation	
use_dycore_barrier	L	.FALSE.		if .TRUE., set an MPI barrier at the beginning of	
				the nonhydrostatic solver (do not use for	
				production runs!)	
itype_exch_barrier	I	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!)	
iorder_sendrecv	I	1		Sequence of send/receive calls:	
				1 = irecv/send	
				$2=\mathrm{isend/recv}$	
				3 = isend/irecv	
				4 = irecv/send with message size blocking	
exch_msgsize	I	8192		Blocking size of exchange messages	$iorder_sendrecv = 4$
itype_comm	I	1		1: use local memory for exchange buffers	
				2: use global memory for exchange buffers	
				3: asynchronous halo communication for dynamical	
				core (NH tria only)	
num_io_procs	I	0		Number of I/O processors (running exclusively for	
				doing I/O)	
pio_type	I	1		Type of parallel I/O. Only used if number of I/O	
				processors greater number of domains.	
				Experimental!	
nh_stepping_thread	s I	1		The number of OpenMP threads to be used by the	
				non-hydrostatic dycore. Only used if the	
				OMP_RADIATION flag is set during	
				compilation. Experimental!	

Parameter	Type	Default	Unit	Description	Scope
radiation_threads	I	1		The number of OpenMP threads to be used by the	
				radiation. Only used if the	
				OMP_RADIATION flag is set during	
				compilation. Experimental!	
use_icon_comm	L	.FALSE.		Enable the use of MPI bulk communication through	
				the icon_comm_lib	
icon_comm_debug	L	.FALSE.		Enable debug mode for the icon_comm_lib	
max_send_recv	I	131072		Size of the send/receive buffers for the	
_buffer_size				icon_comm_lib.	
use_sp_output	L	.FALSE.		Enable this flag if output fields shall be gathered	
				and written in single-precision.	

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

3.5 coupling_nml

Parameter	Type	Default	Unit	Description	Scope
name	С	blank		short name of the coupling field	
dt_coupling	I	0	s	coupling time step / coupling interval	
dt_{model}	I	0	s	model time step	
lag	I	0		offset to coupling event in number of model time	
				steps	
l_time_average	L	.FALSE.		.TRUE.: time averaging between two coupling	
				events	
l_time_accumulatio	n L	.FALSE.		.TRUE.: accumulation of coupling fields in time	
				between two coupling events	
l_diagnostic	L	.FALSE.		.TRUE.: simple diagnostics (min, max, avg) for	
				coupling fields is switched on	
l_activated	L	.FALSE.		.TRUE.: activate the coupling of the respective	
				coupling field	

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

3.6 run_nml

Parameter	Type	Default	Unit	Description	Scope
ldump_states	L	.FALSE.		Dump patch/interpolation/grid refinement state of	
				every patch (after subdivision in case of a parallel	
				run) to a Netcdf file and exit program.	
lrestore_states	L	.FALSE.		Restore patch/interpolation/grid refinement states	
				from NetCDF dump files instead of calculating	
				them.	
dump_filename	С			Filename of dump/restore files, default:	
				" <path>dump_<proc><gridfile>". May contain</gridfile></proc></path>	
				the keyword <path> which will be substituted by</path>	
				model_base_dir, <pre><pre>c> substituted by</pre></pre>	
				"procXofY_", and the grid filename <gridfile>.</gridfile>	
dd_filename	С			Filename of NetCDF domain decomposition dump	
				files, default: " <path>dd_<gridfile>". May</gridfile></path>	
				contain the keyword <path> which will be</path>	
				substituted by model_base_dir, and the grid	
				filename <gridfile>.</gridfile>	
l_one_file_per_pat	chL	.FALSE.		Use one file per patch for all processors.	ldump_states=.TRUE.
				This will decrease the amount of files used for	or
				dump/restore considerably, especially for massively	$lrestore_states=.TRUE.$
				parallel runs on hundreds or thousands of	
				processors.	
				Time for dumping will increase since the file has to	
				be written sequentially, the time for restore should	
				stay roughly the same, however.	

Parameter	Type	Default	Unit	Description	Scope
ldump_dd	L	.FALSE.		Dump the domain decomposition (and a few related	
				fields). This can be done either in a parallel run or	
				in a single-CPU run. When done in a parallel run,	
				the domain decoposition is for the number of	
				parallel processes in use. When done in a	
				single-CPU run, nproc_dd (see below) determines	
				the number of processes for the decomposition.	
				Uses always only one file per patch,	
lread_dd	L	.FALSE.		Read the domain decomposition when dumped with	
				$dump_dd$.	
nproc_dd	I	1		Number of processors for the target domain	dd = TRUE and
				decomposition (only relevant when running on a	a single processor run
				single processor).	
nsteps	I	0		number of time steps of this run.	
dtime	R	600.0	s	time step	
ltestcase	L	.TRUE.		Idealized testcase runs	
ldynamics	L	.TRUE.		Compute adiabatic dynamic tendencies	
iforcing	I	0		Forcing of dynamics and transport by	
				parameterized processes. Use positive indices for	
				the atmosphere and negative indices for the ocean.	
				0: no forcing	
				1: Held-Suarez forcing	
				2: ECHAM forcing	
				3: NWP forcing	
				4: local diabatic forcing without physics	
				5: local diabatic forcing with physics	
				-1: MPIOM forcing (to be done)	
ltransport	L	.FALSE.		Compute large-scale tracer transport	
ntracer	I	0		Number of advected tracers handled by the	
				large-scale transport scheme	
lvert_nest	L	.FALSE.		If set to .true. vertical nesting is switched on (i.e.	
				variable number of vertical levels)	
num_lev	I(max_don	n)31		Number of full levels (atm.) for each domain	lvert_nest=.TRUE.

Parameter	Type	Default	Unit	Description	Scope
nshift	I(max_do	m)0		vertical half level of parent domain which coincides	lvert_nest=.TRUE.
				with upper boundary of the current domain	
ltimer	L	.TRUE.		TRUE: Timer for monitoring thr runtime of specific	
				routines is on $(FALSE = off)$	
timers_level	I	1			
activate_sync_time	rs L	F		TRUE: Timer for monitoring runtime of	
				communication routines $(FALSE = off)$	
msg_level	I	10		controls how much printout is written during	
				runtime.	
				For values less than 5, only the time step is written.	
msg_timestamp	L	.FALSE.		If .TRUE., precede output messages by time stamp.	
test_mode	I	0		Setting a value larger than 0 activates a dummy	iequations = 3
				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 buffer (does not work with	
				nesting and reduced radiation grid because the send	
				buffer may then be empty on some PEs)	
output	L(:)	"vlist","toti	nt"	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;	
				• "vlist" : old, vlist-based output mode;	
				• "nml": new output mode (cf. output_nml);	
				• "totint": computation of total integrals.	
				If the output namelist parameter is not set explicitly, the default setting "vlist", "totint" is assumed.	

$3.7 \quad \mathrm{grid_nml}$

Parameter	Type	Default	Unit	Description	Scope
cell_type	Ι	3		Cell type	
				3: triangular cells	
				4: quadrilateral cells (to be done)	
				6: pentagonal/hexagonal cells	
lplane	L	.FALSE.		planar option	
corio_lat	R	0.0	deg	Center of the f-plane is located at this geographical	lplane=.TRUE.
				latitude	
l_limited_area	L	.FALSE.			
grid_rescale_factor	R	1.0		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.		Specifies if feedback to parent grid is performed.	$n_{dom}>1$
				Setting lfeedback(1)=.false. turns off feedback for	
				all nested domains; to turn off feedback for selected	
				nested domains, set lfeedback(1)=.true. and set	
				".false." for the desired model domains	
$ifeedback_type$	Ι	1		1: incremental feedback	$n_dom > 1$
				2: relaxation-based feedback	
start_time	$R(n_dom)$	0.	s	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	s	Time when a nested domain terminates (namelist	$n_{dom}>1$
				entry is ignored for the global domain)	

Parameter	Type	Default	Unit	Description	Scope
patch_weight	R(n_dom)	0.		If patch_weight is set to a value > 0 for any of the	n_dom>1
				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	L	.FALSE.		If set to .true. is calculated on a reduced grid (=	
				one grid level higher)	
dynamics_grid_	С			Array of the grid filenames to be used by the	
filename				dycore. May contain the keyword <path> which</path>	
				will be substituted by model_base_dir.	
dynamics_parent_	I			Array of the indexes of the parent grid filenames, as	
$\operatorname{grid}_{\operatorname{id}}$				described by the dynamics_grid_filename array.	
				Indexes start at 1, an index of 0 indicates no parent.	
radiation_grid_	С			Array of the grid filenames to be used for the	
filename				radiation model. Filled only if the radiation grid is	
				different from the dycore grid. May contain the	
				keyword <path> which will be substituted by</path>	
				model_base_dir.	
dynamics_radiation	I			Array of the indexes linking the dycore grids, as	
_grid_link				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.	

$3.8 \quad \text{gridref_nml}$

Parameter	Type	Default	Unit	Description	Scope
grf_intmethod_c	I	2		Interpolation method for grid refinement (cell-based	n_dom>1
				dynamical variables):	
				1: parent-to-child copying	
				2: gradient-based interpolation	
$grf_intmethod_ct$	I	2		Interpolation method for grid refinement (cell-based	n_dom>1
				tracer variables):	
				1: parent-to-child copying	
				2: gradient-based interpolation	
$grf_intmethod_e$	I	4		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	
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 variables	$n_{dom}>1$
				(T, p_{sfc}) :	
				1: area-weighted averaging	
				2: bilinear interpolation	
grf_tracfbk	I	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	$n_{dom}>1$
				${\rm edges}\ 3/4$	

Parameter	Type	Default	Unit	Description	Scope
rbf_vec_kern_grf_e	I	1		RBF kernel for grid refinement (edges):	n_dom>1
				1: Gaussian	
				$2: 1/(1+r^2)$	
				3: inverse multiquadric	
rbf_scale_grf_e	R	0.5		RBF scale factor for grid refinement (edges)	n_dom>1
denom_diffu_t	R	135		Deniminator for lateral boundary diffusion of	n_dom>1
				temperature	
denom_diffu_v	R	200		Deniminator for lateral boundary diffusion of	n_dom>1
				velocity	
l_mass_consvcorr	L	.TRUE.		.TRUE.: Apply mass conservation correction in	n_dom>1
				feedback routine	
l_density_nudging	L	.TRUE.		.TRUE.: Apply density nudging near lateral nest	n_dom>1 .AND.
				boundary	led back = .TRUE.

Defined and used in: src/namelists/mo gridref nml.f90

3.9 prepicon nml

Remark: prepicon_nml contains switches controlling the real-data initialization functionality of ICON. There are currently two ways of using it:

- Using the preprocessing tool prep_icon, it is possible to
 - (a) generate the three-dimensional coordinate fields needed by IFS2ICON if IFS2ICON is requested to do the horizontal and vertical interpolation from the IFS grid to the ICON grid
 - (b) convert the hydrostatic set of variables provided by IFS2ICON to the nonhydrostatic set of equations needed by ICONAM, and
 - (c) perform the vertical interpolation to the ICON grid if IFS2ICON is requested to do only the horizontal interpolation step.
- If ICONAM (iequations=3) is combined with NWP physics (iforcing=3), setting ltestcase=.false. activates functionality (c) while running the ICON executable.

Parameter	Type	Default	Unit	Description	Scope
1 di dilicoci	Type	Delaale	01110	Description	Scope

Parameter	Type	Default	Unit	Description	Scope
i_oper_mode	Ι	1		Operating mode if the prep_icon executable is run:	
				1: generate coordinate fields	
				2: convert IFS2ICON output to NH prognostic	
				variables	
				3: do vertical interpolation	
nlev_in	I	91		number of model levels of input data	
nlevsoil_in	I	4		number of soil levels of input data	
zpbl1	R	500.0	m	bottom height (AGL) of layer used for gradient	
				computation	
zpbl2	R	1000.0	m	top height (AGL) of layer used for gradient	
				computation	
l_w_in	L	.FALSE.		Logical switch if vertical wind is provided as input	
l_sfc_in	L	.TRUE.		Logical switch if surface fields are provided as input	
				(mandatory when inwp_surface >0)	
l_zp_out	L	.FALSE.		Logical switch for diagnostic output on pressure	prep_icon only
				and height levels	
l_extdata_out	L	.FALSE.		Logical switch to write extdata fields into output	prep_icon only
l_coarse2fine_mode	L(max_dor	n)FALSE.		If true, apply corrections for coarse-to-fine mesh	
				interpolation to wind and temperature	
ifs2icon_filename	С			Filename of IFS2ICON input file, default	
				" <path>ifs2icon_R<nroot>B<jlev>_DOM<idom>.nc"</idom></jlev></nroot></path>	
				May contain the keywords <path> which will be</path>	
				substituted by model_base_dir, as well as nroot,	
				jlev, and idom defining the current patch.	

Defined and used in: $src/namelists/mo_prepicon_nml.f90$

$3.10 \quad interpol_nml$

Paramete	Default	Unit	Description	Scope

Parameter	Type	Default	Unit	Description	Scope
llsq_lin_consv	L	.FALSE.		conservative (T) or non-conservative (F)	
				least-squares reconstruction for 2nd order (linear)	
				transport	
llsq_high_consv	L	.TRUE.		conservative (T) or non-conservative (F)	
				least-squares reconstruction for high order transport	
lsq_high_ord	I	3		polynomial order for high order reconstruction	
				1: linear	ihadv_tracer=4
				2: quadratic	
				30: cubic (no 3^{rd} order cross deriv.)	
				3: cubic	
rbf_vec_kern_c	I	1		Kernel type for reconstruction at cell centres:	
				1: Gaussian	
				3: inverse multiquadric	
rbf_vec_kern_e	I	3		Kernel type for reconstruction at edges:	
				1: Gaussian	
				3: inverse multiquadric	
rbf_vec_kern_v	I	1		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	
		dependent			
rbf_vec_scale_e	R(n_dom)	resolution-		Scale factor for RBF reconstruction at edges	
		dependent			
rbf_vec_scale_v	R(n_dom)	resolution-		Scale factor for RBF reconstruction at vertices	
		dependent			
nudge_max_coeff	R	0.02		Maximum relaxation coefficient for lateral	
				boundary nudging	
nudge_efold_width	R	2.5		e-folding width (in units of cell rows) for lateral	
				boundary nudging coefficient	
nudge_zone_width	I	8		Total width (in units of cell rows) for lateral	
				boundary nudging zone	
i_cori_method	I	3		Selector for tangential wind reconstruction method	currently only for
					$cell_type=6$

Parameter	Type	Default	Unit	Description	Scope
				1: Almut's method for tangential wind, but PV	
				usage as in TRSK	
				2: method of Thuburn, Ringler, Skamarock and	
				Klemp (TRSK)	
				3: Almut's method for tangential wind and PV	
				usage	
l_corner_vort	L	.TRUE.		switch whether the rhombus averaged corner	i_cori_method=3
				vorticity is averaged to the hexagon (.TRUE.) or	
				the rhombi are directly averaged to the hexagon	
				(.FALSE.)	
l_intp_c2l	L	.FALSE.		If .TRUE. directly interpolate from cell centers to	
				lon-lat points, otherwise do gradient interpolation	
				and reconstruction.	
rbf_dim_c2l	I	10		stencil size for direct lon-lat interpolation: 4 =	
				nearest neighbor, $13 = \text{vertex stencil}$, $10 = \text{edge}$	
				stencil.	
l_mono_c2l	L	.FALSE.		Monotonicity can be enforced by demanding that	
				the interpolated value is not higher or lower than	
				the stencil point values.	

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

$3.11 \quad dynamics_nml$

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

Parameter	Type	Default	Unit	Description	Scope
iequations	I	1		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	

Parameter	Type	Default	Unit	Description	Scope
				-1: hydrostatic ocean	
idiv_method	I	1		Method for divergence computation:	grid_nml:cell_type=3
				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	$idiv_method=2$
sw_ref_height	R	0.9*2.94e4	gm	Reference height of shallow water model used for	
				linearization in the semi-implicit time stepping	
				scheme	
lcoriolis	L	.TRUE.		Coriolis force	

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

$3.12 \quad ha_dyn_nml$

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

Parameter	Type	Default	Unit	Description	Scope
itime_scheme	I	4		Time integration scheme:	
				11: pure advection (no dynamics)	
				12: 2 time level semi implicit (not yet implemented)	
				13: 3 time level explicit	
				14: 3 time level with semi implicit correction	
				15: standard 4th-order Runge-Kutta method	
				(4-stage)	
				16: $SSPRK(5,4)$ scheme (5-stage)	
ileapfrog_startup	I	1		How to integrate the first time step when the	$itime_scheme = 13 \text{ or } 14$
				leapfrog scheme is chosen. $1 = \text{Euler forward}$; $2 = a$	
				series of sub-steps.	
asselin_coeff	R	0.1		Asselin filter coefficient	itime_scheme= 13 or 14
si_2tls	R	0.6		weight of time step $n+1$. Valid range: $[0,1]$	itime_scheme=12

Parameter	Type	Default	Unit	Description	Scope
si_expl_scheme	I	2		scheme for the explicit part used in the 2 time level	itime_scheme=12
				semi-implicit time stepping scheme. $1 = \text{Euler}$	
				forward; $2 = Adams$ -Bashforth 2nd order	
si_cmin	R	30.0	m/s	semi implicit correction is done for eigenmodes with	itime_scheme=14 and
				speeds larger than si_cmin	lsi_3d=.FALSE.
si_coeff	R	1.0		weight of the semi implicit correction	itime_scheme=14
si_offctr	R	0.7			itime_scheme=14
si_rtol	R	1.0e-3		relative tolerance for GMRES solver	itime_scheme=14
lsi_3d	L	.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 state	iequations $\in \{1,2\}$
				when computing the pressure graident force	

3.13 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 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)	iequations=3 and cell_type=3

Parameter	Type	Default	Unit	Description	Scope
rayleigh_type	I	2		Type of Rayleigh damping	cell_type=3
				1: CLASSICAL (requires velocity reference state!)	
				2: Klemp (2008) type	
rayleigh_coeff	R(n_dom)	0.05		Rayleigh damping coefficient $1/\tau_0$ (Klemp, Dudhia,	cell_type=3
				Hassiotis: MWR136, pp.3987-4004)	
damp_height	R(n_dom)	45000	m	Height at which Rayleigh damping of vertical wind	
				starts	
htop_moist_proc	R	22500.0	m	Height above which moist physics and advection of	
				cloud and precipitation variables are turned off	
hbot qvsubstep	R	24000.0	m	Height above which QV is advected with	cell type=3 and
				substepping scheme (must be larger than	ihadv tracer=22 or 32
				htop_moist_proc)	
k2_updamp_coeff	R	2.0e6		enhanced 2nd order diffusion coefficient in upper	cell_type=6,
				damping layer	hdiff order=3
					(Smagorinski)
vwind_offctr	R	0.15		Off-centering in vertical wind solver	cell_type=3
ivctype	I	2		Type of vertical coordinate:	
				1: Gal-Chen hybrid	
				2: SLEVE (uses sleve_nml)	
iadv_rcf	I	4		reduced calling frequency (rcf) for transport	
_				1: no rcf (every dynamics-step)	
				2: transport every 2. step	
				4:	
				Setting odd values (besides 1) requires l nest rcf	
				= .TRUE.	
lhdiff_rcf	L	.TRUE.		.TRUE.: Compute diffusion only at advection time	cell_type=3
_				steps (in this case, divergence damping is applied in	
				the dynamical core)	
divdamp_fac	R	0.004		Scaling factor for divergence damping	$lhdiff_rcf = .TRUE.$
divdamp_order	I	4		Order of divergence damping (2 or 4)	$=$ lhdiff_rcf = .TRUE.
l nest rcf	L	.TRUE.		Synchronize interpolation/feedback calls with	cell type=3
				advection (transport) time steps. l_nest_rcf is	
				automatically reset to .FALSE. if iadv_rcf=1	

Parameter	Type	Default	Unit	Description	Scope
l_masscorr_nest	L	.FALSE.		.TRUE.: Apply mass conservation correction also in	cell_type=3
				nested domain	
iadv_rhotheta	I	2		Advection method for rho and rhotheta:	cell_type=3
				1: centred differences horiz. + vert.	
				2: 2nd order Miura horizontal	
				3: 3rd order Miura horizontal (not recommended)	
igradp_method	I	3		Discretization of horizontal pressure gradient:	cell_type=3
				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	$cell_type=3$.AND.
				diffusion truly horizontally over steep slopes	$hdiff_order=3/5$.AND.
					$lhdiff_temp = .true.$
thslp_zdiffu	R	0.025		Slope threshold above which truly horizontal	$cell_type=3$.AND.
				temperature diffusion is activated	$hdiff_order=3/5$.AND.
					$lhdiff_temp=.true.$
					.AND. $l_zdiffu_t=.true$.
thhgtd_zdiffu	R	200	m	Threshold of height difference between neighboring	cell_type=3 .AND.
				grid points above which truly horizontal	$hdiff_order=3/5$.AND.
				temperature diffusion is activated (alternative	lhdiff_temp=.true.
				criterion to thslp_zdiffu)	.AND. l_zdiffu_t=.true.
exner_expol	R	0.5		Temporal extrapolation (fraction of dt) of Exner	cell_type=3
				function for computation of horizontal pressure	
				gradient	

Parameter	Type	Default	Unit	Description	Scope
l_open_ubc	L	.FALSE.		.TRUE.: Use open upper boundary condition	cell_type=3
				(rather than w=0) to better conserve sea-level	
				pressure in the presence of diabatic heating	
ltheta_up_hori	L	.FALSE.		upstream biased horizontal advection for theta (see	cell_type=6
				also upstr_beta)	
upstr_beta	R	1.0		Selection of order for horiz, theta advection: 3rd	cell_type=6
				order=1.0, 4th order=0.0	
gmres_rtol_nh	R	1.0e-6		relative tolerance for convergence in gmres solver	cell_type=6

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

$3.14 \quad sleve_nml \; (relevant \; if \; nonhydrostatic_nml:ivctype{=}2)$

Parameter	Type	Default	Unit	Description	Scope
min_lay_thckn	R	50	m	Layer thickness of lowermost layer; specifying zero	
				or a negative value leads to constant layer	
				thicknesses determined by top_height and nlev	
top_height	R	23500.0	m	Height of model top	
stretch_fac	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	m	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	L	.FALSE.		read smoothed topography from file (TRUE) or	
				compute internally (FALSE)	

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

$3.15 \quad diffusion_nml$

Parameter	Type	Default	Unit	Description	Scope
lhdiff_temp	L	.TRUE.		Diffusion on the temperature field	
lhdiff_vn	L	.TRUE.		Diffusion on the horizontal wind field	
hdiff_order	I	4 (hydro) 5 (NH)		Order of ∇ operator for diffusion: -1: no diffusion 2: ∇² diffusion (not available for NH model on triangles!) 3: Smagorinsky ∇² diffusion (includes frictional heating for the hexagonal model if lhdiff_temp=.TRUE.) 4: ∇⁴ diffusion 5: Smagorinsky ∇² diffusion combined with ∇⁴ background diffusion as specified via hdiff_efdt_ratio defaults: 2 for hexagonal model, 4 for triangular model; for triangular NH model, 5 is strongly recommended! 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.	24 and 42 currently allowed only in the hydrostatic atm model (run_nml:iequation = 1 or 2).
itype_vn_diffu	I	1		Reconstruction method used for Smagorinsky diffusion: 1: u/v reconstruction at vertices only 2: u/v reconstruction at cells and vertices	iequations=3, hdiff_order=3 or 5
itype_t_diffu	I	1		Discretization of temperature diffusion: 1: $K_h \nabla^2 T$ 2: $\nabla \cdot (K_h \nabla T)$	iequations=3, hdiff_order=3 or 5
k2_pres_max	R	-99.	Pa	Pressure level above which ∇^2 diffusion is applied.	hdiff_order = 24 or 42, and run_nml:iequation = 1 or 2.

Parameter	Type	Default	Unit	Description	Scope
k2_klev_max	I	0		Index of the vertical level till which (from the model	$hdiff_order = 24 \text{ or } 42,$
				top) ∇^2 diffusion is applied. If a positive value is	$and run_nml:iequation =$
				specified for k2_pres_max, k2_klev_max is reset	1 or 2.
				accordingly during the initialization of a model run.	
hdiff_efdt_ratio	R	1.0		ratio of e-folding time to time step (or 2* time step	
				when using a 3 time level time stepping scheme)	
				(only for triangles currently; for triangular NH	
				model, values between 10 and 20 are recommended	
				when using hdiff_order=5)	
hdiff_min_efdt_rati	oR	1.0		minimum value of hdiff_efdt_ratio near model top	iequations=3 .AND.
		(hydro)			$cell_type=3$.AND.
		15.0			$hdiff_order=4$
		(NH)			
hdiff_tv_ratio	R	1.0		Ratio of diffusion coefficients for temperature and	
				normal wind: $T:v_n$	
hdiff_multfac	R	1.0		Multiplication factor of normalized diffusion	n_dom>1
				coefficient for nested domains	
hdiff_smag_fac	R	0.15		Scaling factor for Smagorinsky diffusion	for triangles only with
		(hydro)			iequations=3, for
		0.025			hexagons with
		(NH)			$hdiff_order=3$

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

3.16 io_nml

Parameter	Type	Default	Unit	Description	Scope
out_expname	С	'IHEEEET	TTT'	Outfile basename	
out_filetype	I	2		Type of output format:	
				1: GRIB1 (not yet implemented)	
				2: netCDF	

Parameter	Type	Default	Unit	Description	Scope
lkeep_in_sync	L	.FALSE.		Sync output stream with file on disk after each	
				timestep	
dt_data	R	21600.0	S	Output time interval	
dt_diag	R	86400.		diagnostic integral output interval	
dt_file	R	2592000	S	Time interval of triggering new output file	
dt_checkpoint	R	2592000	S	Time interval for writing restart files. Note that if	
				the value of dt_checkpoint resulting from model	
				default or user's specification is longer than	
				time_nml: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.	
lwrite_vorticity	L	.TRUE.		write out averaged vorticity at vertices	
lwrite_initial	L	.TRUE.		write out initial state	
lwrite_dblprec	L	.FALSE.		write out double precision	
lwrite_oce_timester	p i ng	.FALSE.		write out intermediate ocean vars	
lwrite_divergence	L	.TRUE.		write out divergence at cells	
lwrite_omega	L	.TRUE.		write out vertical velocity in pressure coords.	Always .FALSE. for
					nonhydrostatic and
					shallow water models
lwrite_pres	L	.TRUE.		write out full level pressure	lshallow_water=.FALSE.
lwrite_z3	L	.TRUE.		write out geopotential on full levels	lshallow_water=.FALSE.
lwrite_tracer	L(ntracer)	.TRUE.		write out tracer at cells	
lwrite_tend_phy	L	.TRUE.		Physics induced tendencies.	.TRUE. if
		.FALSE.			iforcing=iecham
		(Scope)			.FALSE. else
lwrite_radiation	L	.FALSE.		Radiation related fields.	Always .FALSE. if
					iforcing=inoforcing,
					iheldsuarez, ildf_dry
lwrite_precip	L	.FALSE.		Precipitation	Always .FALSE. if
					iforcing=inoforcing,
					iheldsuarez, ildf_dry

Parameter	Type	Default	Unit	Description	Scope
lwrite_cloud	L	.FALSE.		Cloud variables	Always .FALSE. if
					iforcing=inoforcing,
					iheldsuarez, ildf_dry
lwrite_tke	L	.TRUE.		TKE	.FALSE.
					Always .FALSE. if
					iforcing=inoforcing,
					iheldsuarez, ildf_dry
lwrite_surface	L	.FALSE.		surface variables	Always .FALSE. if
					iforcing=inoforcing,
					iheldsuarez, ildf_dry
lwrite_extra	L	.FALSE.		debug fields	.TRUE. if inextra_2d
					$/_3d > 0$
					.FALSE. else
inextra_2d	I	0		Number of 2D Fields for diagnostic/debugging	iequations = 3 (to be
				output.	done for $1, 2$)
inextra_3d	I	0		Number of 3D Fields for diagnostic/debugging	iequations $= 3$ (to be
				output.	done for $1, 2$)
lflux_avg	L	.FALSE.		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	
itype_pres_msl	I	1.		Specifies method for computation of mean sea level	
				pressure.	
				1: GME-type extrapolation,	
				2: stepwise analytical integration	

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

$3.17 \quad output_nml$

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.

ılt Unit	Description	Scope
	One of CDI's FILETYPE_XXX constants.	
	/	
4337122		
TWF	,	
	0 11 0	
	ů,	
	The format of this file:	
	One mapping per line, first the internal name, then	
	the name written to NetCDF, separated by an	
	arbitrary number of blanks. The line may also start	
	· ·	
	, , , , , , , , , , , , , , , , , , , ,	
	, v	
	,	
	<u> </u>	
		mode=1
		mode—1
	<u> </u>	
	AWF'	One of CDI's FILETYPE_XXX constants. Possible values: 2 (=FILETYPE_GRB2), 4 (=FILETYPE_NC2), 5 (=FILETYPE_NC4) MWF' 'DWD' - DWD short names (or 'MPIMET', 'CMIP', 'ECMWF') Currently used for setting GRIB2 centre/subcentre information. RJ: For what exactly should that be used? File containing the mapping from internal names to names written to NetCDF. May contain the keyword <path> which will be substituted by model_base_dir. The format of this file: One mapping per line, first the internal name, then the name written to NetCDF, separated by an</path>

Parameter	Type	Default	Unit	Description	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!	
output time unit	I	1		1 = second, 2=minute, 3=hour, 4=day, 5=month,	
				6=year	
output_bounds(3,:)	R	None		post-processing times in units defined by	
				output_time_unit: start, end, increment. There	
				may be specified several triples (up to 100) which	
				must be in increasing order.	
steps_per_file	I	100		Max number of output steps in one output file. If	
				this number is reached, a new output file will be	
				opened.	
include_last	L	.TRUE.		Flag whether to include the last time step	
output_grid	L	.FALSE.		Flag whether grid information is output (in	
				NetCDF output)	
output_filename	С	None		Output filename prefix (which may include path).	
				Domain number, level type, file number and	
				extension will be added, so specifying 'XXX' for	
				output_filename you will end up in a name like	
1		DAT CD		XXX_DOM01_ML_0001.nc	
lwrite_ready	L	.FALSE.		Flag if a "ready file" (sentinel file) should be	
				written at the end of each output stage.	
1 1'		NT.		Not yet implemented.	
ready_directory	С	None		Output directory for ready files.	
1 1: 4()		NT.		Not yet implemented.	
ml_varlist(:)	С	None		Name of model level fields to be output – or 'all'.	
pl_varlist(:)	С	None		Name of pressure level fields to be output – or 'all'.	
hl_varlist(:)	С	None		Name of height level fields to be output – or 'all'.	

Parameter	Type	Default	Unit	Description	Scope
il_varlist(:)	С	None		Name of isentropic level fields to be output – or	
				'all'.	
p_levels(:)	R	None	hPa	pressure levels	
				Not yet implemented.	
				The pressure levels are currently always taken from	
				array plevels in namelist nh_pzlev_nml.	
h_levels(:)	R	None	m	height levels	
ii_icvcis(.)	10	TVOIC	111	Not yet implemented.	
				The height levels are currently always taken from	
				array zlevels in namelist nh pzlev nml.	
i levels(:)	R	None	K	isentropic levels	
		Trone	1	Not yet implemented.	
				The isentropic levels are currently always taken	
				from array ilevels in namelist nh pzlev nml.	
remap	I	0		interpolate horizontally, 0: none, 1: to regular	
T				lat-lon grid, 2: to Gaussian grids, (3:)	
				Currently only 0 and 1 are implemented.	
remap internal	L	.FALSE.		do interpolations online in the model or external	
1 —				(including triggering)	
				Currently unused, interpolations are always done	
				internally.	
$reg_lon_def(3)$	R	None		if remap=1: start, increment, end longitude in	
				degrees	
$reg_lat_def(3)$	R	None		if remap=1: start, increment, end latitude in	
				degrees	
gauss_tgrid_def	I	None		if remap=2: triangular truncation (e.g.63 for T63)	
				for which the Gauss grid should be used	
				Currently unused since Gaussian grids are not	
				implemented.	
$north_pole(2)$	R	0,90		definition of north pole for rotated lon-lat grids.	

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:

group:atmo_ml_vars

group:atmo_pl_vars, group:atmo_zl_vars

group:nh_prog_vars

group:atmo_derived_vars

group:rad_vars
group:precip_vars
group:cloud_diag

group:phys_tendencies

group:land_vars

group:pbl_vars

group:multisnow_vars

basic atmospheric variables on model levels

same set as atmo ml vars, but except pres and height, respectively

additional prognostic variables of the nonhydrostatic model

derived atmospheric variables

tile-averaged variables

10 1 1

Defined and used in: src/namelists/mo name list output.f90

$3.18 \quad meteogram_output_nml$

Parameter	Type	Default	Unit	Description	Scope
lmeteogram_enabled	L(n_dom)	.FALSE.		Flag. True, if meteogram of output variables is	
				desired.	
zprefix	$C(n_dom)$	"METEOG	RAM_"	string with file name prefix for output file	
ldistributed	L(n_dom)	.TRUE.		Flag. Separate files for each PE.	
$n0_{mtgrm}$	I(n_dom)	1		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'			

Defined and used in: src/namelists/mo mtgrm nml.f90

$3.19 \quad nh_pzlev_nml$

Parameter	Type	Default	Unit	Description	Scope
nzlev	I	10		number of height levels	iequations=3
nplev	I	10		number of pressure levels	iequations=3
nilev	I	3		number of isentropes	iequations=3
zlevels	R	10000, 9000, , 1000, 0	m	array of height levels	iequations=3 level ordering from TOA to bottom
plevels	R	100000, 90000, 80000, , 10000	Pa	array of pressure levels	iequations=3 level ordering from TOA to bottom
ilevels	R	340, 320, 300	K	array of isentropic levels	iequations=3 level ordering from TOA to bottom

Defined and used in: $src/namelists/mo_nh_pzlev_nml.f90$

${\bf 3.20 \quad transport_nml \; (used \; if \; run_nml/ltransport=.TRUE.)}$

	Parameter	Type	Default	Unit	Description	Scope
Ī	lvadv_tracer	L	.TRUE.		TRUE : compute vertical tracer advection	
					FALSE: do not compute vertical tracer advection	
Ī	ihadv_tracer	I(ntracer)	2		Tracer specific method to compute horizontal	
					advection:	
			5		0: no horiz. transport	

Parameter	Type	Default	Unit	Description	Scope
				1: upwind (1st order)	
				2: miura (2nd order, lin. reconstr.)	if cell_type=3
				20: miura (2nd order, lin. reconstr.) with	if cell type=3
				subcycling	
				3: miura3 (quadr. or cubic reconstr.)	$lsq_high_ord \in [2,3]$
				22: combination of miura and miura with	if cell type=3
				subcycling	
				32: combination of miura3 and miura with	if cell type=3
				subcycling	
				4: FFSL (quadr. or cubic reconstr.)	lsq high ord $\in [2,3]$
				5: up3 (3rd or 4th order upstream)	if cell_type=6
ivadv tracer	I(ntracer)	3		Tracer specific method to compute vertical	lvadv tracer=TRUE
				advection:	_
				0: no vert. transport	
				1: upwind (1st order)	
				2: $muscl_cfl$ (2nd order, handles $CFL > 1$)	
				20: muscl (2nd order)	
				3: ppm cfl (3^{rd} order, handles CFL > 1)	
				30: ppm (3rd order)	
lstrang	L	.FALSE.		splitting into fractional steps	
				- second order Strang splitting (.TRUE.)	
				- first order Godunov splitting (.FALSE.)	
ctracer_list	С	"		list of tracer names	
itype hlimit	I(ntracer)	3		Type of limiter for horizontal transport:	
		4		0: no limiter	
				3: monotonous flux limiter	ihadv tracer \neq 'iup3[4]'
				4: positive definite flux limiter	
itype_vlimit	I(ntracer)	1		Type of limiter for vertical transport:	
_				0: no limiter	
				1: semi-monotone slope limiter	
				2: monotonous slope limiter	
				4: positive definite flux limiter	

Parameter	Type	Default	Unit	Description	Scope
niter_fct	I	1		number of iterations of monotone flux correction	$ihadv_tracer = 3, 32, 4$
				procedure	$itype_hlimit = 3$
beta_fct	R	1.0		factor for multiplicative spreading of range of	$ihadv_tracer = 3, 32, 4$
				permissible values (limiter)	$itype_hlimit = 3$
				Tentative suggestion: beta_fct=1.0015	
iord_backtraj	I	1		order of backward trajectory calculation:	
				1: first order	
				2: second order (iterative; currently 1 iteration	ihadv_tracer='miura'
				hardcoded)	
igrad_c_miura	I	1		Method for gradient reconstruction at cell center	
				for 2nd order miura	
				1: Least-squares (linear, non-consv)	$ihadv_tracer=2$
				2: Green-Gauss	
ivcfl_max	I	5		determines stability range of vertical PPM-scheme	$ivadv_tracer=3$
				in terms of the maximum allowable CFL-number	
$llsq_svd$	L	.FALSE.		use QR decomposition (FALSE) or SV	
				decomposition (TRUE) for least squares design	
				matrix A	
lclip_tracer	L	.FALSE.		Clipping of negative values	
upstr_beta_adv	R	1.0		parameter to select 3rd order (=1) or 4th order	$ihadv_tracer=iup3$
				(=0) advection, or something inbetween (01)	

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

$3.21 \quad nwp_phy_nml$

Parameter	Type	Default	Unit	Description	Scope
inwp_gscp	I	1		cloud microphysics and precipitation	$run_nml/iforcing = inwp$
				0: none	
				1: hydci (COSMO-EU microphysics)	
				9: Kessler scheme	
qi0	R	0.0	kg/kg	cloud ice threshold for autoconversion	inwp_gscp=1

Parameter	Type	Default	Unit	Description	Scope
qc0	R	0.0	kg/kg	cloud water threshold for autoconversion	inwp_gscp=1
inwp_convection	I	1		convection	$run_nml/iforcing = inwp$
				0: none	
				1: Tiedtke/Bechtold convection	
$inwp_cldcover$	I	3		cloud cover scheme for radiation	$run_nml/iforcing = inwp$
				0: no clouds (only QV)	
				1: grid-scale clouds and QV	
				2: clouds from COSMO turbulence scheme	
				3: clouds from COSMO SGS cloud scheme	
$inwp_radiation$	I	1		radiation	$run_nml/iforcing = inwp$
				0: none	
				1: RRTM radiation	
				2: Ritter-Geleyn radiation	
inwp_satad	I	1		saturation adjustment	$run_nml/iforcing = inwp$
				0: none	
				1:	
inwp_turb	I	1		vertical diffusion and transfer	$run_nml/iforcing = inwp$
				0: none	
				1: COSMO diffusion and transfer	
				2: GME turbulence scheme (to be implemented)	
				3: EDMF-DUALM (work in progress)	
				4: ECHAM diffusion (currently for water only)	
inwp sso	I	1		subgrid scale orographic drag	run nml/iforcing = inwp
				0: none	_ ,
				1: (COSMO) Lott and Miller scheme	
inwp_gwd	I	1		non-orographic gravity wave drag	run nml/iforcing = inwp
1 _0				0: none	_ ,
				1:Orr-Ern-Bechtold-scheme(IFS)	
inwp_surface	I	1		surface scheme	run nml/iforcing = inwp
* _				0: none	_ ,
				1: TERRA	

Parameter	Type	Default	Unit	Description	Scope
ustart_raylfric	R	160.0	m/s	wind speed at which extra Rayleigh friction starts	$inwp_gwd > 0$
efdt_min_raylfric	R	10800.	S	minimum e-folding time of Rayleigh friction	$inwp_gwd > 0$
				(effective for $u > ustart_raylfric + 90 m/s$)	
latm_above_top	L	.FALSE.		.TRUE.: take into account atmosphere above model	$inwp_radiation > 0$
	$(\max_{dom}$)		top for radiation computation	
itype_z0	I	1		Type of roughness length data used for turbulence	$inwp_turb > 0$
				scheme: $1 = \text{including contribution from sub-scale}$	
				orography, $2 = \text{land-cover-related roughness only}$	
dt_conv	R	600.	seconds	time interval of convection call	$run_nml/iforcing = inwp$
	(max_dom)		currently each subdomain has	
				the same value	
$\mathrm{dt}_\mathrm{ccov}$	R	dt_conv	seconds	time interval of cloud cover call	$run_nml/iforcing = inwp$
	$(\max_{dom}$)		currently each subdomain has	
				the same value	
dt_rad	R	1800.	seconds	time interval of radiation call	$run_nml/iforcing = inwp$
	(max_dom)		currently each subdomain has	
				the same value	
dt_sso	R	1200.	seconds	time interval of sso call	$run_nml/iforcing = inwp$
	$(\max_{dom}$)		currently each subdomain has	
				the same value	
dt_gwd	R	1200.	seconds	time interval of gwd call	$ run_nml/iforcing = inwp $
	(max_dom)		currently each subdomain has	
				the same value	

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

$3.22 \quad {\rm radiation_nml}$

Parameter	Type	Default	Unit	Description	Scope
ldiur	L	.TRUE.		switch for solar irradiation:	
				.TRUE.:diurnal cycle,	
				.FALSE.:zonally averaged irradiation	

Parameter	Type	Default	Unit	Description	Scope
nmonth	I	0		0: Earth circles on orbit	
				1-12: Earth orbit position fixed for specified month	
lyr_perp	L	.FALSE.		.FALSE.: transient Earth orbit following VSOP87	
				.TRUE.: Earth orbit of year yr perp of the	
				VSOP87 orbit is perpertuated	
yr_perp	L	-99999		$year used for lyr_perp = .TRUE.$	
isolrad	I	0		Insolation scheme	
				0: Use insolation defined in code.	
				1: Use insolation from external file containing the	
				spectrally resolved insolation averaged over a year	
				(not yet implemented)	
dt_rad	R	7200.	second	time interval of full radiation computation	$run_nml/iforcing =$
					iecham
izenith	I	3		Choice of zenith angle formula for the radiative	
		4 (for		transfer computation.	
		iforcing		0: Sun in zenith everywhere	
		= inwp)		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 ($\sin(\text{time of day}) = 1/\text{pi}$	
				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)	

Parameter	Type	Default	Unit	Description	Scope
irad_h2o	Ι	1		Switches for the concentration of radiative agents	Note: until further notice,
$irad_co2$		2		0: 0.	please use
irad_ch4		3		1: prognostic variable	$irad_h2o = 1$
irad_n2o		3		2: global constant	${ m irad_co2} = 2$
irad_o3		3		3: externally specified	and 0 for all the other
irad_o2		2		irad_aero = 5: Tanre aerosol climatology for	agents for
irad_cfc11		2		$run_nml/iforcing = 3 (NWP)$	${ m run_nml/iforcing} = 2$
irad_cfc12		2		irad_aero = 6: Tegen aerosol climatology for	(ECHAM).
irad_aero		2		$run_nml/iforcing = 3 \text{ (NWP) .AND. itopo } = 1$	
				$irad_o3 = 2$: ozone climatology from MPI	
				$irad_o3 = 4$: 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)	
vmr_co2	R	353.9e-6		Volume mixing ratio of the radiative agents	
vmr_ch4		1693.6e-9			
vmr_n2o		309.5e-9			
vmr_o2		0.20946			
vmr_cfc11		252.8e-12			
vmr_cfc12		466.2e-12			

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

$3.23 \quad nwp_lnd_nml$

Parameter	Type	Default	Unit	Description	Scope
nlev_snow	I	1		number of snow layers	$lmulti_snow=.true.$
				for lmulti_snow=.true.	
ntiles	I	1		number of tiles	

Parameter	Type	Default	Unit	Description	Scope
lsnowtile	L	.FALSE.		.TRUE.: consider snow-covered and snow-free tiles	ntiles>1
				separately	
frlnd_thrhld	R	0.5		fraction threshold for creating a land grid point	ntiles>1
frlndtile_thrhld	R	0.05		fraction threshold for retaining the respective tile	ntiles>1
				for a grid point	
frlake_thrhld	R	0.5		fraction threshold for creating a lake grid point	ntiles>1
frsea_thrhld	R	0.5		fraction threshold for creating a sea grid point	ntiles>1
nztlev	I	2		used time integration scheme	
lmulti_snow	L	.FALSE.		.TRUE. for use of multi-layer snow model	
idiag_snowfrac	I	1		Type of snow-fraction diagnosis: $1 = based$ on SWE	
				only, $2 = \text{more advanced experimental method}$	
lseaice	L	.FALSE.		.TRUE. for use of sea-ice model	
llake	L	.FALSE.		.TRUE. for use of lake model	

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

$3.24 \quad echam_phy_nml$

Parameter	Type	Default	Unit	Description	Scope
lrad	L	.TRUE.		Switch on radiation.	iforcing = 2
lvdiff	L	.TRUE.		Switch on turbulent mixing (i.e. vertical diffusion).	iforcing = 2
lconv	L	.TRUE.		Switch on cumulus convection.	iforcing = 2
lcond	L	.TRUE.		Switch on large scale condensation.	iforcing = 2
lcover	L	.FALSE.		.TRUE. for prognostic cloud cover scheme, .FALSE.	iforcing = 2
				for diagnostic scheme.	Note: $lcover = .TRUE$.
					runs, but has not been
					evaluated (yet) in ICON.
lgw_hines	L	.FALSE.		.TRUE. for atmospheric gravity wave drag by the	iforcing = 2
				Hines scheme	
lssodrag	L	.FALSE.		.TRUE. for subgrid scale orographic drag	iforcing = 2
					Not implemeted yet

Parameter	Type	Default	Unit	Description	Scope
llandsurf	L	.FALSE.		.TRUE. for surface exchanges	iforcing = 2
					Not implemeted yet
lice	L	.FALSE.		.TRUE. for sea-ice temperature calculation	iforcing = 2
					Not implemeted yet
lmeltpond	L	.FALSE.		.TRUE. for calculation of meltponds	iforcing = 2
					Not implemeted yet
lhd	L	.FALSE.		.TRUE. for hydrologic discharge model	iforcing = 2
					Not implemeted yet
lmlo	L	.FALSE.		.TRUE. for mixed layer ocean	iforcing = 2
					Not implemeted yet

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

$3.25 \quad echam_conv_nml$

Parameter	Type	Default	Unit	Description	Scope
iconv	I	1		Choice of cumulus convection scheme.	iforcing = 2 .AND. $lconv$
				1: Nordeng scheme	= .TRUE.
				2: Tiedtke scheme	
				3: hybrid scheme	
ncvmicro	I	0		Choice of convective microphysics scheme.	iforcing = 2 .AND. $lconv$
					= .TRUE.
lmfpen	L	.TRUE.		Switch on penetrative convection.	iforcing = 2 .AND. $lconv$
					= .TRUE.
lmfmid	L	.TRUE.		Switch on midlevel convection.	iforcing = 2 .AND. $lconv$
					= .TRUE.
lmfdd	L	.TRUE.		Switch on cumulus downdraft.	iforcing = 2 .AND. $lconv$
					= .TRUE.
lmfdudv	L	.TRUE.		Switch on cumulus friction.	iforcing = 2 .AND. $lconv$
					= .TRUE.
cmftau	R	10800.		Characteristic convective adjustment time scale.	iforcing = 2 .AND. $lconv$
					= .TRUE.

Parameter	Type	Default	Unit	Description	Scope
cmfctop	R	0.3		Fractional convective mass flux (valid range [0,1])	iforcing = 2 .AND. $lconv$
				across the top of cloud	= .TRUE.
cprcon	R	1.0e-4		Coefficient for determining conversion from cloud	iforcing = 2 .AND. $lconv$
				water to rain.	= .TRUE.
cminbuoy	R	0.025		Minimum excess buoyancy.	iforcing = 2 .AND. $lconv$
					= .TRUE.
entrpen	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. $lconv$
				convective precipitation.	= .TRUE.

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

3.26 vdiff_nml

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

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

$3.27 \quad turbdiff_nml$

Parameter	Type	Default	Unit	Description	Scope
itype_tran	I	2		type of surface-atmosphere transfer	$inwp_turb = 1$
imode_tran	I	1		mode of surface-atmosphere transfer	$inwp_turb = 1$
icldm_tran	I	0		mode of cloud representation in transfer parametr	$inwp_turb = 1$
imode_turb	I	3		mode of turbulent diffusion parametrization	$inwp_turb = 1$
icldm_turb	I	2		mode of cloud representation in turbulence	$inwp_turb = 1$
				parametr	
itype sher	I	1		type of shear production for TKE	inwp turb = 1

Parameter	Type	Default	Unit	Description	Scope
ltkesso	L	.FALSE.		calculation SSO-wake turbulence production for	$inwp_turb = 1$
				TKE	
ltkecon	L	.FALSE.		consider convective buoyancy production for TKE	$inwp_turb = 1$
lexpcor	L	.FALSE.		explicit corrections of the implicit calculated turbul.	$inwp_turb = 1$
				diff.	
ltmpcor	L	.FALSE.		consideration of thermal TKE-sources in the	$inwp_turb = 1$
				enthalpy budget	
lprfcor	L	.FALSE.		using the profile values of the lowest main level	$inwp_turb = 1$
				instead of the mean value of the lowest layer for	
				surface flux calulations	
lnonloc	L	.FALSE.		nonlocal calculation of vertical gradients used for	$inwp_turb = 1$
				turbul. diff.	
lcpfluc	L	.FALSE.		consideration of fluctuations of the heat capacity of	$inwp_turb = 1$
				air	
limpltkediff	L	.TRUE.		consideration of fluctuations of the heat capacity of	$\mathrm{inwp_turb} = 1$
				air	
itype_wcld	I	2		type of water cloud diagnosis	$inwp_turb = 1$
itype_synd	I	2		type of diagnostics of synoptical near surface	$inwp_turb = 1$
				variables	
lconst_z0	L	.FALSE.		TRUE: horizontally homogeneous roughness lenght	$inwp_turb = 1$
				z0	
const_z0	R	0.001	m	value for horizontally homogeneous roughness	$inwp_turb = 1$
				lenght z0	lconst_z0=.TRUE.

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

$3.28 \quad gw_hines_nml \; (Scope: \; lgw_hines = .TRUE. \; in \; echam_phy_nml)$

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

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

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

4 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.

$4.1 \quad ha_testcase_nml \; (Scope: \; ltestcase=.TRUE. \; and \; iequations=[0,1,2] \; in \; run_nml)$

Parameter	Type	Default	Unit	Description	Scope
ctest_name	С	'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.

Parameter	Type	Default	Unit	Description	Scope
				'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 physics	$lshallow_water=.FALSE.$
					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 wave	$lshallow_water=.FALSE.$
				'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.
$rotate_axis_deg$	R	0.0	\deg	Earth's rotation axis pitch angle	ctest_name= 'Will_2',
					'Will_3', 'JWs', 'JWw',
	D	0.01	4 /		'PA', 'DF1234'
gw_brunt_vais	R	0.01	1/s	Brunt Vaisala frequency	ctest_name= 'GW'
gw_u0	R	0.0	m/s	zonal wind parameter	ctest_name= 'GW'
gw_lon_deg	R	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	m/s	amplitude of the wave pertubation	ctest_name= 'JWw'
, , 1 1	D	00.0	(?)		23.50337(0);
mountctr_lon_deg	R	90.0	deg	longitude of mountain peak	$ctest_name = 'MRW(2)'$

Parameter	Type	Default	Unit	Description	Scope
mountctr_lat_deg	R	30.0	deg	latitude of mountain peak	ctest_name= 'MRW(2)'
mountctr_height	R	2000.0	m	mountain height	ctest_name= 'MRW(2)'
mountctr_half_widt	hR	1500000.0	m	mountain half width	ctest_name= 'MRW(2)'
mount_u0	R	20.0	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 test.	ctest_name= 'HS'
				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 the	ctest_name= 'HS'
				Held-Suarez test.	
hs_vn_ptb_scale	R	1.	m/s	Magnitude of the random noise added to the initial	ctest_name= 'HS'
				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'
ape_sst_case	С	'sst1'		SST distribution selection	ctest name='APE'
1				'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.	
		l .		<u>'</u>	

Parameter	Type	Default	Unit	Description	Scope
ildf_init_type	I	0		Choice of initial condition for the Local diabatic	$ctest_name = 'LDF'$
				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	L	.TRUE.		Shape of local diabatic forcing:	ctest_name=
				.TRUE.: local diabatic forcing symmetric about the	'LDF','LDF-Moist'
				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

$4.2 \quad nh_testcase_nml \; (Scope: \; ltestcase=.TRUE. \; and \; iequations=3 \; in \; run_nml)$

Parameter	Type	Default	Unit	Description	Scope
nh_test_name	С	'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.	
				'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.	

Parameter	Type	Default	Unit	Description	Scope
				'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 nh': Initializes the APE experiments. With	
				the jabw test case, including moisture.	
				'wk82': Initializes the Weisman Klemp test case	l limited area =.TRUE.
				'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_rest_200': atmosphere at rest test	lcoriolis = .FALSE.
				(Schaer-type mountain)	
				'dcmip_mw_2x': nonhydrostatic mountain	lcoriolis = .FALSE.
				waves triggered by Schaer-type mountain	
				'dcmip_gw_31': nonhydrostatic gravity waves	
				triggered by a localized perturbation (nonlinear)	
				'dcmip_gw_32': nonhydrostatic gravity waves	$l_{limited_area} = .TRUE.$
				triggered by a localized perturbation (linear)	and lcoriolis $=$.FALSE.
				'dcmip_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	
jw_up	R	1.0	m/s	amplitude of the u-perturbation in jabw test case	nh_test_name='jabw'

Parameter	Type	Default	Unit	Description	Scope
u0_mrw	R	20.0	m/s	wind speed for mrw(2) and mwbr_const cases	$nh_test_name =$
					$'mrw(2)_nh'$ and
					$'mwbr_const'$
mount_height_mrw	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	R	1500000.0	m	half width of mountain in mrw(2), mwbr_const	$nh_test_name =$
				and bell	$'mrw(2)_nh',$
					'mwbr_const' and 'bell'
mount_lonctr_mrw_	_dReg	90.	degrees	lon of mountain center in mrw(2) and mwbr_const	$nh_test_name =$
					$'mrw(2)_nh'$ and
					'mwbr_const'
mount_latctr_mrw_	d a g	30.	degrees	lat of mountain center in mrw(2) and mwbr_const	$nh_test_name =$
					$'mrw(2)_nh'$ and
					'mwbr_const'
temp_i_mwbr_cons	t R	288.0	K	temp at isothermal lower layer for mwbr_const case	$nh_test_name =$
					'mwbr_const'
p_int_mwbr_const	R	70000.	Pa	pres at the interface of the two layers for	$nh_test_name =$
				mwbr_const case	'mwbr_const'
bruntvais_u_mwbr_	cRnst	0.025	1/s	constant brunt vaissala frequency at upper layer for	$nh_test_name =$
				mwbr_const case	'mwbr_const'
mount_height	R	100.0	m	peak height of mountain	nh_test_name= 'bell'
layer_thickness	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	Ι	2		level number for which the layer is still flat and not	$layer_thickness>0$
1 0	D	0.0	/	terrain-following	1 1 21 212
nh_u0	R	0.0	m/s	initial constant zonal wind speed	nh_test_name = 'bell'
nh_t0	R	300.0	K	initial temperature at lowest level	nh_test_name = 'bell'
nh_brunt_vais	R	0.01	1/s	initial Brunt-Vaisala frequency	$nh_test_name = 'bell'$

Parameter	Type	Default	Unit	Description	Scope
torus_domain_lengt	hR	100000.0	m	length of slice domain	nh_test_name = 'bell',
					lplane=.TRUE.
rotate_axis_deg	R	0.0	deg	Earth's rotation axis pitch angle	nh_test_name= 'PA'
lhs nh vn ptb	L	.TRUE.		Add random noise to the initial wind field in the	nh test name=
				Held-Suarez test.	'HS_nh'
lhs_fric_heat	L	.FALSE.		add frictional heating from Rayleigh friction in the	nh_test_name=
				Held-Suarez test.	'HS_nh'
hs_nh_vn_ptb_sca	leR	1.	m/s	Magnitude of the random noise added to the initial	nh_test_name=
				wind field in the Held-Suarez test.	'HS_nh'
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'
ape_sst_case	С	'sst1'		SST distribution selection	nh_test_name='APE_nh'
				'sst1': Control experiment	
				'sst2': Peaked experiment	
				'sst3': Flat experiment	
				'sst4': Control-5N experiment	
				'sst_qobs': Qobs SST distribution exp.	
$linit_tracer_fv$	L	.TRUE.		Finite volume initialization for tracer fields	pure advection tests, only
lcoupled_rho	L	.FALSE.		Integrate density equation 'offline'	pure advection tests, only
qv_max_wk	R	0.014	Kg/kg	maximum specific humidity near	nh_test_name='wk82'
				the surface, range 0.012 - 0.016	
				used to vary the buoyancy	
u_infty_wk	R	20.	m/s	zonal wind at infinity height	nh_test_name='wk82'
				range 0 45.	
				used to vary the wind shear	
bub_amp	R	2.	K	maximum amplitud of the thermal perturbation	nh_test_name='wk82'
bubctr_lat	R	0.	deg	latitude of the center of the thermal perturbation	nh_test_name='wk82'
bubctr_lon	R	90.	deg	longitude of the center of the thermal perturbation	nh_test_name='wk82'
bubctr_z	R	1400.	m	height of the center of the thermal perturbation	nh_test_name='wk82'
bub_hor_width	R	10000.	m	horizontal radius of the thermal perturbation	nh_test_name='wk82'
bub_ver_width	R	1400.	m	vertical radius of the thermal perturbation	nh_test_name='wk82'

Parameter	Type	Default	Unit	Description	Scope
itype_atmo_ana	Ι	1		kind of atmospheric profile:	nh_test_name=
	ı			1 piecewise N constant layers	'g_lim_area'
	1			2 piecewise polytropic layers	
itype_anaprof_uv	Ι	1		kind of wind profile:	nh_test_name=
	ı			1 piecewise linear wind layers	'g_lim_area'
	ı			2 constant zonal wind	
	ı			3 constant meridional wind	
itype_topo_ana	I	1		kind of orography:	$nh_test_name =$
	ı			1 schaer test case mountain	'g_lim_area'
	ı			2 gaussian_2d mountain	
	ı			3 gaussian_3d mountain	
	ı			any other no orography	
nlayers_nconst	Ι	1		Number of the desired layers with a constant	$nh_test_name =$
	ı			Brunt-Vaisala-frequency	'g_lim_area' and
	ı				itype_atmo_ana=1
p_base_nconst	R	100000.	Pa	pressure at the base of the first N constant layer	$nh_test_name =$
	ı				'g_lim_area' and
	1				itype_atmo_ana=1
theta0_base_nconst	R	288.	K	potential temperature at the base of the first N	$nh_test_name =$
	ı			constant layer	'g_lim_area' and
	ı				itype_atmo_ana=1
h_nconst	R(nlayers_	n00,n\$5)00.,	m	height of the base of each of the N constant layers	$nh_test_name =$
	ı	12000.			'g_lim_area' and
	ı				itype_atmo_ana=1
N_nconst	R(nlayers_	noonst)	1/s	Brunt-Vaisala-frequency at each of the N constant	nh_test_name=
	1			layers	'g_lim_area' and
	1				itype_atmo_ana=1
rh_nconst	R(nlayers_	n 0 ofast)	%	relative humidity at the base of each N constant	nh_test_name=
	1			layers	'g_lim_area' and
					itype_atmo_ana=1
rhgr_nconst	R(nlayers_	n0onst)	%	relative humidity gradient at each of the N constant	nh_test_name=
	1			layers	'g_lim_area' and
	1				itype_atmo_ana=1

Parameter	Type	Default	Unit	Description	Scope
nlayers_poly	I	2		Number of the desired layers with constant gradient	nh_test_name=
				temperature	'g_lim_area' and
					$itype_atmo_ana=2$
p_base_poly	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_	p 6 ly)12000.	m	height of the base of each of the polytropic layers	nh_test_name=
					'g_lim_area' and
					itype_atmo_ana=2
t_poly	R(nlayers_	p 28 8., 213.	K	temperature at the base of each of the polytropic	$nh_test_name =$
				layers	'g_lim_area' and
					$itype_atmo_ana=2$
rh_poly	R(nlayers_	p 6l §) 0.2	%	relative humidity at the base of each of the	nh_test_name=
				polytropic layers	'g_lim_area' and
					itype_atmo_ana=2
rhgr_poly	R(nlayers_	p ōly -)5, 0.	%	relative humidity gradient at each of the polytropic	$nh_test_name =$
				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
h_linwind	R(nlayers_	li 0wi250 0.	m	height of the base of each of the linear wind layers	$nh_test_name =$
					'g_lim_area' and
					itype_anaprof_uv=1
u_linwind	R(nlayers_	li 5 w 10 d)	m/s	zonal wind at the base of each of the linear wind	nh_test_name=
				layers	'g_lim_area' and
					itype_anaprof_uv=1
ugr_linwind	R(nlayers_	lionwiond)	1/s	zonal wind gradient at each of the linear wind layers	nh_test_name=
					'g_lim_area' and
					itype_anaprof_uv=1
vel_const	R	20.	m/s	constant zonal/meridional wind	nh_test_name=
				(itype_anaprof_uv=2,3)	'g_lim_area' and
					$itype_anaprof_uv=2,3$

Parameter	Type	Default	Unit	Description	Scope
mount_lonc_deg	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	R	250.	m	h0 parameter for the schaer mountain	nh_test_name=
					'g_lim_area' and
					itype_topo_ana=1
schaer_a	R	5000.	m	-a- parameter for the schaer mountain,	nh_test_name=
				also half width in the north and south side of the	'g_lim_area' and
				finite ridge to round the sharp edges	itype_topo_ana=1,2
schaer_lambda	R	4000.	m	lambda parameter for the schaer mountain	nh_test_name=
					'g_lim_area' and
					itype_topo_ana=1
lshear_dcmip	L	FALSE		run dcmip_mw_2x with/without vertical wind	nh_test_name=
				shear	'dcmip_mw_2x'
				FALSE: dcmip_mw_21: non-sheared	
				TRUE : dcmip_mw_22: sheared	
halfwidth_2d	R	10000.	m	half length of the finite ridge in the north-south	nh_test_name=
				direction	'g_lim_area' and
					itype_topo_ana=1,2
m_height	R	1000.	m	height of the mountain	nh_test_name=
					'g_lim_area' and
					itype_topo_ana=2,3
m_width_x	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	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

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

5 External data

$5.1 \quad extpar_nml \; (Scope: \; itopo=1 \; in \; run_nml)$

Parameter	Type	Default	Unit	Description	Scope
itopo	Ι	0		0: analytical topography/ext. data	
				1: topography/ext. data read from file	
n_iter_smooth_top	o I(n_dom)	0		iterations of topography smoother	itopo = 1
fac_smooth_topo	R	0.015625		pre-factor of topography smoother	$n_{iter_smooth_topo} >$
					0
heightdiff_threshold	R(n_dom)	3000.	m	height difference between neighboring grid points	
				above which additional local nabla2 diffusion is	
				applied	
l_emiss	L	.TRUE.		read and use external surface emissivity map	itopo = 1
extpar_filename	С			Filename of external parameter input file, default:	
				" <path>extpar_<gridfile>". May contain the</gridfile></path>	
				keyword <path> which will be substituted by</path>	
				model_base_dir.	

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

6 External packages

$6.1 \quad art_nml$

Parameter	Type	Default	Unit	Description	Scope
lart	L	.FALSE.		main switch for ART-package	
lemi_volc	L	.FALSE.		Emission of volcanic ash	
lconv tracer	L	.FALSE.		Convection of tracers	
lwash tracer	L	.FALSE.		Washout of tracers	
lrad volc	L	.FALSE.		Radiative impact of volcanic ash	
lcld tracer	L	.FALSE.		Impact on clouds	

Defined and used in: src/namelists/mo art nml.f90

7 Information on vertical level distribution

The hydrostatic and nonhydrostatic models need hybrid vertical level information to generate the terrain following coordinates. The hybrid level specification is stored in <icon home>/hyb_params/HYB_PARAMS_<nlev>. The hydrostatic model assumes to get pressure based coordinates, the nonhydrostatic model expects height based coordinates. For further information see <icon home>/hyb params/README.