Working with the ICON Model

Practical Exercises
for NWP Mode and ICON-ART

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0. Preface

The ICON (ICOsdedral Nonhydrostatic) modeling framework (Zängl et al., 2015) is a joint project between the Deutscher Wetterdienst (DWD) and the Max-Planck-Institute for Meteorology (MPI-M) for developing a unified next-generation global numerical weather prediction (NWP) and climate modeling system.

The main goals formulated in the initial phase of the collaboration are

- better conservation properties than in the existing global models, with the obligatory requirement of exact local mass conservation and mass-consistent transport,
- better scalability on future massively parallel high-performance computing architectures,
- the availability of some means of static mesh refinement. ICON is capable of mixing one-way nested and two-way nested grids within one model application, combined with an option for vertical nesting. This allows the global grid to extend into the mesosphere (which facilitates the assimilation of satellite data) whereas the nested domains extend only into the lower stratosphere in order to save computing time.
- applicability on a wide range of scales down to \( O(1 \text{ km}) \) and beyond (which of course requires a nonhydrostatic dynamical core).

The ICON modeling framework became operational in DWD’s forecast system in January 2015. During the first six months only global simulations were executed with a horizontal grid spacing of 13 km and 90 vertical levels. Starting from July 21st, 2015, model simulations have been complemented by a nesting region over Europe. Finally, in January 2018, the global 40 member ICON-EPS (Ensemble Prediction System) was released for the operational service at DWD.

The model source code has been made available for scientific use under an institutional license since 2015.

0.1. How This Document Is Organized

Not all topics in this manuscript are covered during the workshop. Therefore, the manuscript can be used as a textbook, similar to a user manual for the ICON model. Readers are assumed to have a basic knowledge of the design and usage of numerical weather prediction models.

Even though the chapters in this textbook are largely independent, they should preferably not be treated in an arbitrary order.
• For getting started with the ICON model: read Chapters 1 – 5.
• New users who are interested in the regional model should read Chapter 6 in addition.
• More advanced topics are covered by Chapters 7 – 11.

Paragraphs describing common pitfalls and containing details for advanced users are marked by the symbol 🛠.

At the end of the document a number of exercises is provided (see page 215). These exercises revisit the topics of the individual chapters and range from easy tests to the setup of complex forecast simulations.

To some extent this document can also be used as a reference manual. We refer to the index on page 255 for a quick look-up of namelist parameters.

0.2. How to Obtain a Copy of the ICON Model Code

The ICON model is distributed under an institutional license\(^1\). To obtain a grant of license that must be signed and returned to the DWD, please contact icon@dwd.de or follow the information on the public ICON web site

https://code.mpimet.mpg.de/projects/iconpublic

Additionally, we have established the mailing list icon-community@mpimet.mpg.de to stay in touch with the ICON community. Please visit https://listserv.gwdg.de/mailman/listinfo/icon-community and subscribe to this list in order to receive announcements about new releases and features.

Data Services

On the ICON web page under the mentioned URL you will also find the access to the grid generator web service (see Section 2.1.4), and the web links to ICON’s official grid download site and GRIB2 definitions.

DWD has made a number of model forecast data sets publicly available, mostly free of charge (with a retention of 48 h). This service has started in July 2017 and can be reached under https://opendata.dwd.de/weather/icon. See the content description under https://www.dwd.de/EN/ourservices/opendata/opendata.html for a list of spatial data sets.

For further data requests with respect to DWD operational data products please contact datenservice@dwd.de.

\(^1\) An individual licensing procedure has not yet been released by April 2018.
0.3. Further Documentation

The ICON model is accompanied by various other manuals and documentation. An extensive list is available and constantly kept up-to-date in the documentation section of the public ICON web site

https://code.mpimet.mpg.de/projects/iconpublic

We restrict ourselves to a small subset in the following.

Scientific Documentation.

Up to now there is no comprehensive scientific documentation available. In this respect, we refer to the publication Zängl et al. (2015) and the references cited therein.

Recent information on ICON’s hydrostatic dynamical core and the LES model can be found in Wan et al. (2013), Dipankar et al. (2015), Heinze et al. (2017).

Detailed information and evaluation of the atmospheric component of ICON using the climate physics package is given by Giorgetta et al. (2018), Crueger et al. (2018).

The extended modules for Aerosols and Reactive Trace gases (ART) are described in Rieger et al. (2015), Schröter et al. (2018).

A description of the ocean component ICON-O within the ICON modeling system (which is not covered by this tutorial) can be found in Korn (2017), Korn and Danilov (2017).

Technical Documentation.

For model users who intend to process data products of DWD’s operational runs, the DWD database documentation may be a valuable resource. It can be found (in English language) on the DWD web site


A complete list of namelist switches can be found in the namelist documentation

icon/doc/Namelist_overview.pdf

which is deployed together with the code.

The pre- and post-processing tools of the DWD ICON Tools collection are described in more detail in the DWD ICON Tools manual, see Prill (2014).

Finally, please note the FAQ section on the ICON web site which covers a variety of common pitfalls.
1. Installation of the ICON Model Package

The purpose of this tutorial is to give you some practical experience in installing and running the ICON model package. Exercises are carried out on the supercomputers at DWD but the principal steps of the installation can directly be transferred to other systems.

1.1. The ICON Model Package

The source code for the ICON model package consists of the following three components:

- **The ICOsahedral Nonhydrostatic model (ICON)**
  The ICON code that is used for this tutorial has been derived from the ICON release v2.3.00 with additional enhancements for ICON-LAM (state January 2019). It is close to DWD’s currently operational icon-nwp version (see below for explanation). The code also contains the ocean model developed at MPI-M which is, however, not covered by this tutorial.

- **ICON-ART for aerosols and reactive trace gases**
  The ART module, where ART stands for Aerosols and Reactive Trace gases, is an extension of the ICON model to enable the simulation of gases, aerosol particles and related feedback processes in the atmosphere. The module is provided by the Karlsruhe Institute of Technology (KIT) and requires a separate license (see Section 10.1).

- **DWD ICON Tools**
  The ICON Tools are a set of command-line tools for remapping, extracting and querying ICON data files. They are based on a common library and written in Fortran 90/95 and Fortran 2003.

**ICON-NWP code version:** The versioning of ICON is a bit complex and reflects the parallel development in several “flavors” like “atmosphere”, “ocean”, and several more. There are four important branches tagging versions that reached certain milestones: icon-aes (atmosphere in the Earth system, mainly echam physics in the atmosphere), icon-nwp (numerical weather prediction, mainly dynamics and physics of the LEM and NWP configurations of the atmospheric model), icon-oes (ocean in the Earth system), icon-les (land in the Earth system). The common release version integrates the stable components of all branches.

Each tag contains all model components, but the latest tag of icon-aes may not contain the most recent developments of the ocean physics although these are already included into the latest icon-oes tag.
1.1.1. Directory Layout

Figure 1.1 shows a brief description of the directory structure of the ICON model and of the directories containing the test case data under the root tree.

```
icon_tutorial
  └── documentation
  └── reference
    ├── test_cases
      └── case_idealized
      └── case_realdata
      └── case_lam
    └── icon
      └── src
        └── art
      └── support
      └── config
      └── include
        └── lapack
        └── blas
        └── externals
        └── build
      └── dwd_icon_tools
```

e.g. tutorial hand-outs

solutions to exercises, example output

test case 1: idealized experiment *(see Ch. 4)*

test case 2: real-data experiment *(see Ch. 5)*

test case 3: limited area experiment *(see Ch. 6)*

Fortran sources

ART code *(see Ch. 10)*

C99 support library and utility routines

platform configuration, see Section 1.2.1

C library interface

numerical utility library

numerical utility library

external submodules *(calendar etc.)*

build directory with sources and binary

utilities, e.g. for remapping data

Figure 1.1.: Directory structure of the ICON model and of the directories containing the test case data under the root tree. Note that the ICON-ART source code modules are distributed in a separate tarball.

The ICON model code is located in the directory `icon`. The most important subdirectories are described in the following:

**Subdirectory build**

Within the `build` directory, a subdirectory with the name of your computer architecture is created during compilation. When you open this newly created folder you find a `bin` subdirectory containing the ICON binary `icon` and several other subdirectories containing the compiled module files.
1.1 The ICON Model Package

Subdirectory config

Inside the config directory, different machine-dependent configurations are stored in configuration script files (see Section 1.2.1).

Subdirectory src

Within the src directory we have the source code of ICON including the main program and ICON modules. The modules are organized in several subdirectories:

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

The configuration of ICON run-time settings is implemented within the modules inside src/configure_model and src/namelists. Modules regarding the configuration of idealized test cases can be found inside src/testcases.

The dynamics of ICON are implemented inside src/atm_dyn_iconam and the physical parameterizations inside src/atm_phy_nwp. Surface parameterizations can be found inside src/lnd_phy_nwp.

Shared infrastructure modules for 3D and 4D variables are located within src/shared. Routines that are primarily related to horizontal grids and 2D fields (e.g. external parameters) are stored within src/shr_horizontal.

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

Input and output modules are stored in src/io.

The ICON code comes with its own LAPACK and BLAS sources. For performance reasons, these libraries may be replaced by machine-dependent optimizations. However, please note that LAPACK and BLAS routines are not actively used by the nonhydrostatic model.

1.1.2. Libraries Needed for Data Input and Output

Especially for I/O tasks, the ICON model package requires external libraries. Two data formats are implemented in the package to read and write data from or to disk: GRIB and NetCDF.

- GRIB (GRIdded Binary) is a standard defined by the World Meteorological Organization (WMO) for the exchange of processed data in the form of grid point values expressed in binary form. GRIB coded data consists of a continuous bit-stream made of a sequence of octets (1 octet = 8 bits). Please note that the ICON model does support only the GRIB2 version of the standard.

- NetCDF (Network Common Data Form) is a set of software libraries and machine-independent data formats that support the creation, access, and sharing of array-oriented scientific data. NetCDF files contain the complete information about the dependent variables, the history, and the fields themselves. The NetCDF file format is also used for the definition of the computational mesh (grid topology).

For more information on NetCDF see http://www.unidata.ucar.edu.
To work with the formats described above the following libraries are utilized by the ICON model package. For this training course, the paths to access these libraries on the used computer system are already specified in the Makefile.

**The Climate Data Interfaces (CDI) – support/cdilib.c**

This library has been developed and implemented by the Max-Planck-Institute for Meteorology in Hamburg. It provides a C and Fortran interface to access climate and NWP model data. Among others, supported data formats are GRIB1/2 and NetCDF. A condensed copy of the CDI is distributed together with the ICON model package. Note that the CDI are also used by the DWD ICON Tools.

For more information see [https://code.mpimet.mpg.de/projects/cdi](https://code.mpimet.mpg.de/projects/cdi).

**The ECMWF ecCodes package\(^1\) – libecodes.a, libecodes_f90.a**

The European Centre for Medium-Range Weather Forecasts (ECMWF) has developed an application programmers interface (API) to pack and unpack GRIB1 as well as GRIB2 formatted data. For reading and setting meta-data, the ecCodes package uses the so-called key/value approach, which means that all the information contained in the GRIB message is retrieved through alphanumeric names. Indirect use of this ecCodes library in the ICON model is implemented through the CDI.

In addition to the GRIB library, there are some command-line tools to provide an easy way to check and manipulate GRIB data from the shell. Amongst them, the most important ones are `grib ls` and `grib dump` for listing the contents of a grib file, and `grib set` for (re)-setting specific key/value pairs.

For more information on ecCodes we refer to the ECMWF web page:

[https://confluence.ecmwf.int/display/ECC](https://confluence.ecmwf.int/display/ECC)

---

**Installation:** The source code for the ecCodes package can be downloaded from the ECMWF web page.

Please refer to the README for installing the ecCodes libraries, which is done with a configure script. Check the following settings:

- The ecCodes package can make use of optional JPEG packing of the GRIB records, but this requires the installation of additional libraries. Since the ICON model does not apply this packing algorithm, the support for JPEG can be disabled during the configure step with the option `--disable-jpeg`.

---

\(^1\)ecCodes is an evolution of the former GRIB-API software package. To facilitate the alternative use of both libraries, ICON implements only backward-compatible API function names. The GRIB-API libraries, however, would be `libgrib_api.a, libgrib_api_f90.a`.
1. Installation

1.1 The ICON Model Package

- To use statically linked libraries and binaries you should set the configure option `--enable-shared=no`.

```
./configure --prefix=/your/install/dir \
--disable-jpeg --enable-shared=no
```

After the configuration has finished, the ecCodes library can be built with `make` and then `make install`.

### GRIB Definition Files

An installation of the ecCodes package always consists of two parts: First, there is the binary compiled library itself with its functions for accessing GRIB files. But, second, there is the definitions directory which contains plain-text descriptions of meta data.

GRIB definition files are external text files which constitute a kind of parameter database. They describe the decoding rules and the keys which are used to identify the meteorological fields. For example, these definition files contain information about the variable short name and the corresponding GRIB code triplet.

The DWD definition files for the ecCodes package can be obtained via

https://opendata.dwd.de/weather/lib/grib/

The new directory needs to be communicated to the ecCodes package at run-time by setting the `GRIB_DEFINITION_PATH` environment variable:

```
export
GRIB_DEFINITION_PATH=/your/path/definitions.edzw:/your/path/definitions
```

Here, the definitions directory provided by ECMWF is extended by DWD’s own installation (`definitions.edzw`). Note that both paths have to be specified in this environment variable, and that `definitions.edzw` has to be the first! The current setting of the definition files path can be displayed with the command-line tool `codes_info`.

In contrast to the GRIB triplet, the short name, e.g., “T” for temperature, is not stored in data files. The definition file therefore constitutes an essential link: If the definition files in two institutes are different from each other it is possible that the same data file shows the record “OMEGA” on one site (our DWD system), while the same GRIB record bears the short name “w” on the other site (both have the same GRIB triplet `discipline=0,parameterCategory=2,parameterNumber=8`).

The ICON model accesses its input data by their name (“shortName” key). Therefore the DWD-specific definition files (“EDZW”=DWD Offenbach) are essential for the read-in process. In theory, the above situation could be solved by changing all field names in the ICON name list setup, where possible. However, it is likely that further related errors may follow in the ICON model when this searches for a specific variable name. In this case you might need to change the definition files after all.
Note that for writing GRIB2 files, the ICON model does not use DWD-specific shortNames. Therefore, the model output can be written in GRIB2 format without the proper definition files at hand.

**The NetCDF library – libnetcdf.a**

A special library, the NetCDF library, is necessary to write and read data using the NetCDF format. This library also contains tools for manipulating and visualizing the data (*ncdump* utility, see Section 9.1.1).

If the library is not yet installed on your system, you can get the source code and documentation from

http://www.unidata.ucar.edu/software/netcdf/index.html

This includes a description how to install the library on different platforms. Please make sure that the F90 package is also installed, since the model reads and writes grid data through the F90 NetCDF functions.

Note that there exists a restriction regarding the file size. While the classic NetCDF format could not deal with files larger than 2 GiB the new NetCDF-4/HDF5 format permits storing files as large as the underlying file system supports. However, NetCDF-4/HDF5 files are unreadable to the NetCDF library before version 4.0.

### 1.2. Configuring and Compiling the Model Code

This section explains the configuration process of the ICON model. It is assumed that the libraries and programs discussed in Section 1.1.2 are present on your computer system.

For convenience, the compiler version and the ecCodes version are documented in the log output of each model run.

### 1.2.1. Computer Platforms

For a small number of HPC platforms settings are provided with the code, for example

**Cray XC 40 cluster** (*xc.dwd.de*)

432 compute nodes Intel Haswell (24 cores/node, 64 GiB memory)
864 compute nodes Intel Broadwell (36 cores/node, 64 GiB memory)

- **MPI:** Cray MPICH 7.4.3
- **NetCDF:** Version 4.3.2
- **Compiler:** Cray Fortran v8.4.1

1Currently position # 278 on Top500 list November 2018.
1.2 Configuring and Compiling the Model Code

<table>
<thead>
<tr>
<th>Fortran Compiler</th>
<th>Working Version</th>
</tr>
</thead>
<tbody>
<tr>
<td>GNU</td>
<td>gcc v6.4.0</td>
</tr>
<tr>
<td>Cray</td>
<td>ftn v8.4.1</td>
</tr>
<tr>
<td>Intel</td>
<td>ifort v17.0.6</td>
</tr>
<tr>
<td>NAG</td>
<td>nagfor v6.0.1064</td>
</tr>
</tbody>
</table>

Table 1.1.: Compiler versions which are known to successfully build the ICON code (state January 2019)

HLRE-3 cluster\(^2\) “Mistral“ (DKRZ Hamburg)
1550 compute nodes Intel Haswell (2 CPUs/node, 12 cores/CPU)
1750 compute nodes Intel Broadwell (2 CPUs/node, 18 cores/CPU)

- MPI: Intel MPI library 2017.0.098
- NetCDF: Version 4.4.2
- Compiler: GNU compiler gcc 6.2.0

Both compiler setups are defined in the configuration file config/mh-linux.

Other architecture-dependent settings may be specified in the directory icon/config – in most cases in the file config/mh-linux therein. To add a specific compiler or change your compiler flags, you have to enter the config/mh-OS according to your operating system OS. Be warned that you need some knowledge about Unix / Linux, compilers and Makefiles to make the necessary adjustments w.r.t. the computing environment.

Due to the usage of modern Fortran 2003/2008 features, ICON places high demands on the compilers. Please make also sure that a compatible compiler for the C99 routines in the package is available. Both components, the Fortran parts and the C parts use a source preprocessor. Table 1.1 provides a list of compilers which are known to successfully build the recent ICON code. There’s a good chance that more recent compiler versions might work as well. However, be aware that this is not necessarily the case. E.g. many of the newer versions of the Cray compiler (ftn v8.4.1 < version < ftn 8.6.0) might have issues.

**Intel ifort compiler:** When compiling with ifort before version 17.0.1, the default behavior is for the compiler not to use the Fortran rules for automatic allocation on intrinsic assignment. You will need to use an option like -assume realloc-lhs.

The ICON model supports different modes of parallel execution, see Section 7.3 for details:

- In the first place, ICON has been implemented for distributed memory parallel computers using the Message Passing Interface (MPI). MPI is a library specification, proposed as a standard by a broadly based committee of vendors, implementors, and users, see [http://www.mcs.anl.gov/research/projects/mpi](http://www.mcs.anl.gov/research/projects/mpi).

\(^2\)Currently position # 64 on Top500 list November 2018.
Moreover, on multi-core platforms, the ICON model can run in parallel using shared-memory parallelism with OpenMP. The OpenMP API is a portable, scalable technique 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 multi-core systems and shared-memory systems, see http://openmp.org.

Finally, note that although ICON has been implemented for distributed memory parallel computers using the Message Passing Interface (MPI), the model can also be installed on sequential computers, where MPI and/or OpenMP are not available. Of course, this execution mode limits the model to rather small problem sizes.

### 1.2.2. Configuring and Compiling

**Pre-compiled Binaries:** Users of the Cray XC 40 system may find recent pre-compiled binaries in the following subdirectory:

```
~routfor/routfox/abs
```

Users of the ECMWF system may find recent pre-compiled binaries in the following subdirectory:

```
/sc1/home/zde/routfox/abs
```

A configure file is provided that takes over the main work of generating the compilation setup. This `autoconf` configuration is used to analyze the computer architecture (hardware and software) and sets user specified preferences, e.g. the compiler. As explained in Section 1.2.1 these preferences are read from `config/mh-<OS>`, where `<OS>` is the identified operating system.

To configure the source code, please log into the Cray XC 40 login node `xce` and change into the subdirectory `icon`. Then type:

```
./configure --with-fortran=compiler
```

where `compiler` is `{gcc,nag,intel,pgi,cray}`. The default is `gcc`. Here, for the DWD platform, please choose the option `--with-fortran=cray`. If the configuration process fails, take a look at the text file `config.log` that is created during the configuration process. This technical log file may contain hints on which particular library has been found missing.

With the Unix command `make` the programs are compiled and all object files are linked to create the binaries. On most machines you can also compile the routines in parallel by using the GNU-make with the command `gmake -j np`, where `np` gives the number of processors to use (`np` typically about 8).

During the compilation process, a subdirectory with the name of your computer architecture is created within the `build` directory. In this subdirectory, a `bin` subdirectory containing the binary `icon` and several further subdirectories containing the compiled module
files can be found. If problems occur during the linking process, check the prerequisites that have been outlined in Section 1.2.1.

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
```

---

**Note for advanced users:** Only the Cray XC 40 platform does not require an explicit "--with-openmp" option for hybrid parallel binaries. If one uses, e.g., the Intel Fortran compiler, then this option is explicitly needed in the configure process.

## 1.3. The DWD ICON Tools

### 1.3.1. General Overview

The DWD ICON Tools provide a number of utilities for the pre- and post-processing of ICON model runs. All of these tools can run in parallel on multi-core systems (OpenMP) and some offer an MPI-parallel execution mode in addition. We give a short overview over several tools in the following and refer to the documentation Prill (2014) for details.

**ICONGRIDGEN**  
*Used in Section 2.1.3*

The `icongridgen` tool is a simple grid generator. It creates icosahedral grids from scratch, which can be fed into the ICON model. Alternatively, an existing global or local grid file is taken as input and parts of this input grid (or the whole grid) are refined via bisection. No storage of global grids is necessary and the tool also provides an HTML plot of the grid.

**ICONREMAP**  
*Used in Sections 2.2.3, 2.3*

The `iconremap` utility is especially important for pre-processing the initial data for the basic test setups in this manuscript. `iconremap` (ICOsaheiral NONhydrostatic model REMAPping) is a utility program for horizontally interpolating ICON data onto regular grids and vice versa. Besides, it offers the possibility to interpolate between triangular grids of different grid spacing.

The `iconremap` tool reads and writes data files in GRIB2 or NetCDF file format. For triangular grids an additional grid file in NetCDF format must be provided.
Several interpolation algorithms are available: Nearest-neighbor remapping, radial basis function (RBF) approximation of scalar fields, area-weighted formula for scalar fields, RBF interpolation for wind fields from cell-centered zonal, meridional wind components $u$, $v$ to normal and tangential wind components at edge midpoints of ICON triangular grids (and reverse), and barycentric interpolation.

Note that \texttt{iconremap} only performs a horizontal remapping, while the vertical interpolation onto the model levels of ICON is handled by the model itself.

\textbf{ICONSUB} \hspace{1cm} – \textit{Used in Section 2.3}

The \texttt{iconsub} tool (ICOsahedral Nonhydrostatic model \textit{SUB}grid extraction) allows “cutting” sub-areas out of ICON data sets.

After reading a data set on an unstructured ICON grid in GRIB2 or NetCDF file format, the tool comprises the following functionality: It may ‘cut out” a subset, specified by two corners and a rotation pole (similar to the COSMO model). Alternatively, a boundary region of a local ICON grid, specified by parent-child relations, may be extracted. This execution mode is especially important for the setup of the limited area model ICON-LAM.

Multiple sub-areas can be extracted in a single run of \texttt{iconsub}. Finally, the extracted data is stored in GRIB2- or NetCDF file format.

\textbf{ICONGPI}

The \texttt{icongpi} tool (ICOsahedral Nonhydrostatic model \textit{Grid Point Information}) is a utility program for searching / accessing individual grid points of an ICON grid. It can be used to determine cells in a triangular grid corresponding to a given geographical position and to determine the geographical position for a given cell index.

\subsection*{1.3.2. Configuring and Compiling the DWD ICON Tools}

To compile the DWD ICON Tools binaries, log into the Cray XC 40 login node \texttt{xce} and change into the subdirectory \texttt{icontools} by typing

\begin{verbatim}
cd dwd_icon_tools/icontools/
\end{verbatim}

You get a list of available compile targets by typing \texttt{make}. The following output is exemplary and may differ from your current version:

\begin{verbatim}
DWD ICONTOOLS
A set of command-line tools for remapping, extracting and querying
\end{verbatim}
1.3 The DWD ICON Tools

ICON data files.

Available Makefile targets:

<table>
<thead>
<tr>
<th>target name</th>
<th>platform</th>
<th>parallelization</th>
<th>compiler</th>
</tr>
</thead>
<tbody>
<tr>
<td>local</td>
<td>local DWD workstation,</td>
<td>OpenMP</td>
<td>gfortran</td>
</tr>
<tr>
<td>local_mpi</td>
<td>local DWD workstation,</td>
<td>OpenMP + MPI</td>
<td></td>
</tr>
<tr>
<td>ibmp7_mpi</td>
<td>IBM Power 7,</td>
<td>OpenMP + MPI</td>
<td>XLF</td>
</tr>
<tr>
<td>cray_mpi</td>
<td>Cray XE 6 / Cray XC 40,</td>
<td>OpenMP + MPI</td>
<td>Cray FTN</td>
</tr>
<tr>
<td>lce_intel</td>
<td>lce</td>
<td>OpenMP</td>
<td>Intel</td>
</tr>
<tr>
<td>lce_intel_dbg</td>
<td>lce with debugging flags</td>
<td>OpenMP</td>
<td>Intel</td>
</tr>
<tr>
<td>lce_intel_mpi</td>
<td>lce</td>
<td>OpenMP + MPI</td>
<td>Intel</td>
</tr>
<tr>
<td>lce_intel_mpi_dbg</td>
<td>lce with debugging flags</td>
<td>OpenMP + MPI</td>
<td>Intel</td>
</tr>
<tr>
<td>cca_intel_mpi</td>
<td>cca ECMWF</td>
<td>OpenMP + MPI</td>
<td>Intel</td>
</tr>
<tr>
<td>fh2_intel_mpi</td>
<td>FH2 KIT</td>
<td>OpenMP + MPI</td>
<td>Intel</td>
</tr>
<tr>
<td>ucl_intel_mpi</td>
<td>UC1 KIT</td>
<td>OpenMP + MPI</td>
<td>Intel</td>
</tr>
<tr>
<td>knl_intel</td>
<td>DWD KNL</td>
<td>OpenMP</td>
<td>Intel</td>
</tr>
<tr>
<td>nag_mpi</td>
<td>Mistral DKRZ (experimental)</td>
<td>OpenMP + MPI</td>
<td>nagfor</td>
</tr>
<tr>
<td>gcc_mpi</td>
<td>Mistral DKRZ (experimental)</td>
<td>OpenMP + MPI</td>
<td>gfortran</td>
</tr>
<tr>
<td>mistral_intel</td>
<td>Mistral DKRZ (production)</td>
<td>OpenMP + MPI</td>
<td>intel</td>
</tr>
<tr>
<td>gfortran_openmpi_generic</td>
<td>Generic target, adapt paths! OpenMP + OpenMPI</td>
<td>gfortran</td>
<td></td>
</tr>
<tr>
<td>clean</td>
<td>remove all object files, libraries and executables</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For example, the binary for the Cray XC 40 can be created by typing

```
make cray_mpi
```

\textit{ICON Tools Libraries:} The DWD ICON Tools are divided into several independent libraries which can be linked against user applications.

First, there is a low-level API (interpolation, load grid, query point etc.) where all the necessary data is provided via subroutine interfaces. Second, there exists an over-arching \texttt{libicontools.a} sub-library which is used by the high-level API (\texttt{iconremap}, \texttt{iconsut}, \texttt{icongpi}, \texttt{icongridgen}) and handles namelists.

The DWD ICON utilities use the ecCodes package for reading data in GRIB2 format. The ecCodes package is indirectly accessed by the Climate Data Interface (CDI).
1. Installation
2. Necessary Input Data

Before anything else, preparation is the key to success.

Alexander Graham Bell

Besides the source code of the ICON package and the technical libraries, several data files are needed to perform runs of the ICON model. There are four categories of necessary data: Horizontal grid files, external parameters, and data describing the initial state (DWD analysis or ECMWF IFS data). Finally, running ICON in limited-area mode in addition requires accurate boundary conditions sampled at regular time intervals.

2.1. Horizontal Grids

In order to run ICON, it is necessary to load the horizontal grid information as an input parameter. This information is stored within so-called grid files. For an ICON run, at least one global grid is required. For model runs with nested domains, additional grids are necessary. Optionally, a reduced radiation grid for the global domain may be used (see Section 3.10).

The following nomenclature has been established: In general, by $R_nB_k$ we denote a grid that originates from an icosahedron whose edges have been initially divided into $n$ parts, followed by $k$ subsequent edge bisections. See Figure 2.1 for an illustration of the grid creation process. The total number of cells in a global ICON grid $R_nB_k$ is given by $n_{\text{cells}} := 20 n^2 4^k$. The cell circumcenters serve as data sites for most ICON variables. As an exception, the orthogonal normal wind is given at the midpoints of the triangle edges and is measured orthogonal to the edges.

The effective mesh size can be estimated as

$$\Delta x \approx \frac{5050}{(n 2^k)} \text{[km]}.$$  \hspace{1cm} (2.1)

This formula is motivated as follows:

The average triangle area is calculated from the surface of the Earth divided by the number of triangles of the grid. Then, we imagine a square with the same area and define its edge length as the effective mesh size of the triangular grid. This results in

$$\Delta x = \sqrt{\frac{4 R_{\text{earth}}^2 \pi}{n_{\text{cells}}}} = \frac{R_{\text{earth}}}{n 2^k} \sqrt{\frac{\pi}{5}} \approx \frac{5050}{(n 2^k)} \text{[km]},$$
2. Input Data

Figure 2.1.: Illustration of the grid construction procedure. The original spherical icosahedron is shown in red, denoted as $R_{1B00}$ following the nomenclature described in the text. In this example, the initial division ($n=2$; black dotted), followed by one subsequent edge bisection ($k=1$) yields an $R_{2B01}$ grid (solid lines).

where $S_{\text{earth}}$ and $R_{\text{earth}}$ define the Earth’s surface and radius, respectively.

Note that by construction, each vertex of a global grid is adjacent to exactly 6 triangular cells, with the exception of the original vertices of the icosahedron, the pentagon points, which are adjacent to only 5 cells.

Dual Hexagonal Grid

The centers of the equilateral triangles contained in each triangle of the original icosahedron after triangulation are defined by the intersection of the angle bisectors (which are at the same time also altitudes) of the triangle. The centers of the triangles form a hexagonal grid that is called to be dual to the grid of triangle vertices. On the ICON grid, the centers of the slightly distorted triangles form a dual grid of slightly distorted hexagons.

Spring Dynamics Optimization

This grid on the sphere will be optimized in a next step by so-called spring dynamics. We give the idea of the optimization only and refer to Tomita et al. (2002) for an in-depth description of the algorithm.

Imagine that we have a collection of springs all of them of the same strength and length. First, we attach a mass to each triangle vertex and fix it with glue on the circumscribed sphere. We replace each edge by one of the springs. Depending on the actual length of the
edge, we have to tension some springs a bit more for the larger triangles, less for smaller ones. Now, the glue is melted away and the vertices move until an equilibrium is reached provided that there is some friction of the mass points on the sphere.

By this procedure, we will obtain a slightly different grid of triangles which are still slightly distorted and of unequal size, however, the vertices reached positions that reflect some “energy minimum”. These triangles are the basis of the ICON horizontal grid. Such a grid has particularly advantageous numeric properties. The North and South Pole of the Earth are chosen to be located at two vertices of the icosahedron that are opposite to each other.

**ICON “Nests”**

ICON has the capability for running

- global simulations on a single grid,
- limited area simulations (see Chapter 6), and
- global or limited-area simulations with refined nests (so called patches or domains).

For the subtle difference between nested and limited-area setups the reader is referred to Section 6.1. Section 3.9.1 explains the exchange of information between domains.

Additional topological information is required for ICON’s refined nests: Each “parent” triangle is split into four “child” cells. In the grid file only child-to-parent relations are stored while the parent-to-child relations are computed in the model setup. The local numbering of the four child cells (see Fig. 2.2) is also computed in the model setup.

The refinement information may be provided in a separate file (suffix `-grfinfo.nc`), which happens to be the case especially for legacy data sets. This optional grid connectivity file acts as a fallback at model startup if the expected information is not found in the main grid file.

Finally, note that the data points on the triangular grid are the cell circumcenters. Therefore the global grid data points are located closely to nest data points, but they do not coincide exactly.

**2.1.1. ICON Grid Files**

The unstructured triangular ICON grid resulting from the grid generation process is represented in NetCDF format. This file stores coordinates and topological index relations between cells, edges and vertices.

The most important data entries of the main grid file are

- **cell** (INTEGER dimension)
  number of (triangular) cells
- **vertex** (INTEGER dimension)
  number of triangle vertices
2. Input Data

Figure 2.2.: Illustration of the parent-child relationship in refined grids. *Left:* Triangle subdivision and local cell indices. *Right:* The grids fulfill the ICON requirement of a right-handed coordinate system \([\vec{e}_t, \vec{e}_n, \vec{e}_w]\). Note: the primal tangent and the dual cell normal are aligned but do not necessarily coincide!

- **edge** (INTEGER dimension)
  number of triangle edges

- **clon, clat** (double array, dimension: \#triangles, given in radians)
  longitude/latitude of the midpoints of triangle circumcenters

- **vlon, vlat** (double array, dimension: \#triangle vertices, given in radians)
  longitude/latitude of the triangle vertices

- **elon, elat** (double array, dimension: \#triangle edges, given in radians)
  longitude/latitude of the edge midpoints

- **cell_area** (double array, dimension: \#triangles)
  triangle areas

- **vertex_of_cell** (INTEGER array, dimensions: [3, \#triangles])
  The indices vertex_of_cell(:,i) denote the vertices that belong to the triangle i. The vertex_of_cell index array is ordered counter-clockwise for each cell.

- **edge_of_cell** (INTEGER array, dimensions: [3, \#triangles])
  The indices edge_of_cell(:,i) denote the edges that belong to the triangle i.

- **clon/clat_vertices** (double array, dimensions: [\#triangles, 3], given in radians)
  clon/clat_vertices(1,:) contains the longitudes/latitudes of the vertices that belong to the triangle i.

- **neighbor_cell_index** (INTEGER array, dimensions: [3, \#triangles])
  The indices neighbor_cell_index(:,i) denote the cells that are adjacent to the triangle i.

- **zonal/meridional_normal_primal_edge**: (INTEGER array, \#triangle edges)
  components of the normal vector at the triangle edge midpoints.

- **zonal/meridional_normal_dual_edge**: (INTEGER array, \#triangle edges)
  These arrays contain the components of the normal vector at the facets of the dual control volume.
  Note that each facet corresponds to a triangle edge and that the dual normal matches the direction of the primal tangent vector but signs can be different.
2.1 Horizontal Grids

2.1.2. Download of Predefined Grids

For fixed domain sizes and resolutions a list of grid files has been pre-built for the ICON model together with the corresponding reduced radiation grids and the external parameters.

The contents of the primary storage directory are regularly mirrored to a public web site for download, see Figure 2.3 for a screenshot of the ICON grid file server. The download server can be accessed via

http://icon-downloads.mpimet.mpg.de

The pre-defined grids are identified by a centre number, a subcentre number and a numberOfGridUsed, the latter being simply an integer number, increased by one with every new grid that is registered in the download list. Also contained in the download list is a tree-like illustration which provides information on parent-child relationships between global and local grids, and global and radiation grids, respectively.

Note that the grid information of some of the older grids (no. 23 – 40) is split over two files: The users need to download the main grid file itself and a grid connectivity file (suffix -grfinfo.nc).

2.1.3. Grid Generator: Invocation from the Command Line

There are (at least) three grid generation tools available for the ICON model: One grid generation tool has been developed at the Max-Planck-Institute for Meteorology by L. Linardakis. Second, in former releases, the ICON model itself was shipped together with a standalone tool grid_command. This program has finally been replaced by another grid generator which is contained in the DWD ICON Tools.

1See the repository https://code.mpimet.mpg.de/projects/icon-grid-generator.
In this section we will discuss the grid generator `icongridgen` that is contained in the DWD ICON Tools, because this utility also acts as the backend for the publicly available web tool. The latter is shortly described in Section 2.1.4. It is important to note, however, that this grid generator is not capable of generating non-spherical geometries like torus grids.

**Minimum version required:** Grid files that have been generated by the `icongridgen` tool contain only child-to-parent relations while the parent-to-child relations are computed in the model setup. Therefore these grids only work with ICON versions newer than ~ September 2016.

### Grid Generator Namelist Settings

The DWD ICON Tools utility `icongridgen` is mainly controlled using a Fortran namelist. The command-line option that is used to provide the name of this file and other available settings are summarized via typing

```
icongridgen --help
```

The Fortran namelist `gridgen.nml` contains the filename of the parent grid which is to be refined and the grid specification is set for each child domain independently. For example (COSMO-EU nest) the settings are

```
dom(1)%region_type = 3
dom(1)%lrotate = .true.
dom(1)%hwidth_lon = 20.75
dom(1)%hwidth_lat = 20.50
dom(1)%center_lon = 2.75
dom(1)%center_lat = 0.50
dom(1)%pole_lon = -170.00
dom(1)%pole_lat = 40.00
```

For a complete list of available namelist parameters we refer to the documentation (Prill (2014)).

The `icongridgen` grid generator checks for overlap with concurrent refinement regions, i.e. no cells are refined which are neighbors or neighbors-of-neighbors (more precisely: vertex-neighbor cells) of parent cells of another grid nest on the same refinement level. Grid cells which violate this distance rule are “cut out” from the refinement region. Thus, there is at least one triangle between concurrent regions.

**Minimum distance between child nest boundary and parent boundary:** A second, less well-known constraint sometimes leads to unexpected (or even empty) result grids: In the case that the parent grid itself is a bounded regional grid, no cells can be refined that are part of the indexing region (of width `bdy_indexing_depth`) in the vicinity of the parent grid’s boundary.
2.1 Horizontal Grids

Settings for ICON-LAM

When the grid generator `icongridgen` is targeted at a limited area setup (for ICON-LAM), two important namelist settings must be considered:

- **Identifying the grid boundary zone.** In Section 2.3 we will describe how to drive the ICON limited area model. Creating the appropriate boundary data makes the identification of a sufficiently large boundary zone necessary.
  
  This indexing is enabled through the following namelist setting in `gridgen_nml`:
  
  ```
  bdy_indexing_depth = 14.
  ```
  
  This means that 14 cell rows starting from the nest boundary are marked and can be identified in the ICON-LAM setup, which is described in Section 2.3. See Fig. 2.10 for an illustration of such a boundary zone.

- **Generation of a coarse-resolution radiation grid (see Section 3.10 for details).**
  
  The creation of a separate (local) parent grid with suffix `*.parent.nc` is enabled through the following namelist setting in `gridgen_nml`:
  
  ```
  dom(:,%lwrite_parent = .TRUE.
  ```
  
  Note that a grid whose child-to-parent indices are occupied by such a coarse grid can no longer be used in a standard feedback-loop together with a global grid.

2.1.4. Grid Generator: Invocation via the Web Interface

A web service has been made available to help users with the generation of custom grid files. After entering grid coordinates through an online form, this web service creates a corresponding ICON grid file together with the necessary external parameter file.

You will need to log in via the user `icon-web`. For the necessary login password – or if you have trouble accessing the web service – please contact `icon@dwd.de`. Then, visit the web page of the grid generator

```
https://webservice.dwd.de/cgi-bin/spp1167/webservice.cgi
```

The web form is more or less self-explanatory. The settings reflect the namelist parameters of the `icongridgen` grid generator tool that runs as the first stage of the web service. These are explained in the following. The second stage, the ExtPar tool, does not require further settings (with the exception of the surface tiles setting, see below).

The tool is capable of generating multiple grid files at once. **Please note that the web-based grid generation submits a batch job to DWD’s HPC system and takes some time for processing!** Due to limited computing resources a threshold is imposed: the maximum grid size which can be generated is 3 000 000 cells. Of course, larger grid sizes are allowed when using the grid generator apart from the the web interface.

Finally all results (and log files) are packed together into a `*.tar.gz` archive and the user is informed via e-mail about its FTP download site. Additionally, a web browser visualization of the grids based on OpenStreetMap is provided, see Fig. 2.4.
2. Input Data

**ICON Model Tutorial**

**Figure 2.4.** Web browser screenshot of the web-based ICON grid generator tool.

**Step 1: Choosing the Base Grid**

The web-based generation of ICON grids and their corresponding external parameter (ExtPar) data sets starts from an “input file”, which can be chosen from a pull-down menu with a pre-defined list of grids. These grids are identical to those of the download list described in Section 2.1.2.

It is also possible to start from a ”synthetic” base grid $R_nB_k$ by specifying $n$ and $k$, following the algorithm by Sadourny et al. (1968), see p. 17.

Note that when starting from an already existing file, the base grid will not be modified by the grid generator, and it will not be stored together with the generator output. The user merely chooses a sub-region on the globe where the base grid is extracted and refined. The result grid will then have half of the grid size of the base grid.

**Step 2: Specify Global Options**

A number of options will be applied to all produced data sets (“global options”):

- **“centre”, “subcentre”**
  These numbers are stored in the output meta-data section for the identification of the originating / generating sub-center. The values are defined by the WMO, e.g. DWD: 78/0 (see WMO’s Common Code Table C-11 for additional values).
• "spring dynamics optimization"
  The ICON grids are based on the spherical icosahedron, but they are post-processed by an iterative algorithm inspired by elastostatics, see the explanation on “spring dynamics optimization” in Section 2.1.

  – "max. number of iterations"
    maximum number of pseudo-time stepping steps of the elastostatics model.

  – "beta factor"
    stiffness coefficient of the elastostatics model.

  – "fixed lateral boundary"
    If this checkbox is set, then the boundary vertices of regional grids are not moved during the optimization. Thus they will still coincide with vertices of the base grid.

  Recommended: Do not alter the default settings unless you know the details of the underlying algorithm.

• "ExtPar: Tile Mode"
  The web-based generator can produce ExtPar data with and without additional information for surface tiles (see the explanation on p. 50). The data files do not differ in the number of fields, but only in the way some fields are defined near coastal regions.

  The – recommended – default has the tile data enabled. This setting can be changed by disabling the corresponding checkbox in the HTML form.

Step 3: Sub-domain Name and Parent Grid ID

By default, the web form contains only the input fields to specify a single grid. However, more domain specifications can be added to the generator by a click on the "Add another domain" button (or removed by clicking "Remove latest domain").

Numbering: In the web form the base grid is denoted by "#1" while the created domains are denoted by numbers "#2", "#3" and so on.

In the simplest case the domains specify multiple nests on the same refined level. In this case, the "parent grid ID" is always set to "1" (base grid).

Besides, domains can be nested, for example

  Germany → Europe → Global.

Then, the grid for Germany has "parent grid ID=2", and the Europe grid has "parent grid ID=1".

The currently chosen hierarchy of grids is graphically depicted in the top right corner of the web form.

• "domain name"
  Each domain requires a name (string) which will be used as a name prefix for the resulting files.
• "number of grid used"
  This setting will be written into the grid meta-data. It is part of a GRIB2 mechan-
  ism to link data files to their underlying grid files, see also the explanation in
  Section 2.1.2. For regional domains which are not used operationally, we suggest to
  choose arbitrary but distinguishable integer numbers.

• "write parent grid"
  Enable this checkbox when your grid file is to be used with ICON-LAM and a reduced
  radiation grid. Note that in this case the grid cannot be used as a nest in standard
  (non-LAM) mode – see the corresponding remark in Section 2.1.3 above.

Step 4: Specify the Grid Type/Shape

• "global"
  In this case, all cells of the base grid are refined, resulting in a grid with \((4N)\)
  triangles if the base grids consists of \(N\) triangles. No further settings are required to
  specify this grid.

• "rectangular"
  This specifies a sub-region to be refined by a center latitude/longitude (in degrees)
  and a size of \(2 \times \)"half height" for the latitude and \(2 \times \)"half width" in terms of the
  longitude.

  – "rotate" / "north-pole"
    By default, the latitude-longitude coordinates for the rectangular refinement
    area are based on the standard North Pole 90N0E. You may use, however, a
    rotated pole similar to the grids of the COSMO model.

• "circular"
  This defines a circular-shaped refinement region with a given center and radius (in
  degrees).

Finally, having filled out all necessary fields of the web form, click on the “Proceed” button.
The grid generation job is inserted into DWD’s processing queue and you will be informed
via e-mail about its completion.

2.1.5. Which Grid File is Related to My Simulation Data?

ICON data files do not (completely) contain the description of the underlying grid. This
is an important consequence of the fact that ICON uses unstructured, pre-generated com-
putational meshes which are the result of a relatively complex grid generation process.
Therefore, given a particular data file, one question naturally arises: Which grid file is
related to my simulation data?

The answer to this question can be obtained with the help of two meta-data items which
are part of every ICON data and grid file (either a NetCDF global file attribute or a
GRIB2 meta-data key):
2.2 Initial Conditions

- **numberOfGridUsed**
  This is simply an integer number, as explained in the previous sections. The `numberOfGridUsed` helps to identify the grid file in the public download list. If the `numberOfGridUsed` differs between two given data files, then these are not based on the same grid file.

- **uuidOfHGrid**
  This acronym stands for universally unique identifier and corresponds to a binary data tag with a length of 128 bits. The UUID can be viewed as a fingerprint of the underlying grid. Even though this is usually displayed as a hexadecimal number string, the UUID identifier is not human-readable. Nevertheless, two different UUIDs can be tested for equality or inequality.

The meta-data values for `numberOfGridUsed` and `uuidOfHGrid` offer a way to track the underlying grid file through all transformations in the scientific workflow, for example in:

- external parameter files
- analysis data for forecast input
- data files containing the diagnostic output
- checkpointing files (restarting).

### 2.2. Initial Conditions

Global numerical weather prediction (NWP) is an initial value problem. The ability to make a skillful forecast heavily depends on the accuracy with which the present atmospheric (and surface/soil) state is known. Running forecasts with a limited area model in addition requires accurate boundary conditions sampled at regular time intervals.

Initial conditions are usually generated by a process called *data assimilation*. Data assimilation combines irregularly distributed (in space and time) observations with a short term forecast of a general circulation model (e.g. ICON) to provide a "best estimate" of the current atmospheric state. Such *analysis products* are provided by several global NWP centers. In the following we will present and discuss the data sets that can be used to drive the ICON model.

Each computational domain, i.e. also a nested sub-region, requires a separate initial data file. A "workaround" to start a nested simulation without the need to provide initial data for the nest is discussed in Section 5.2. The basic preprocessing steps for ICON are visualized in Figure 2.5.

### 2.2.1. Obtaining DWD Initial Data

The most straightforward way to initialize ICON is to make use of DWD’s analysis products, which are generated operationally every 3 hours. They are available in GRIB2 format. Note that for so-called *idealized test cases* no initial conditions must be read in. All necessary state variables are preset by analytical values.
2. Input Data

Driving Model / Data Assimilation

Figure 2.5.: Basic preprocessing steps for ICON (without limited area mode) which include the generation of grids and external parameters as well as the remapping of initial conditions. The grid generation process is described in Sections 2.1 – 2.4. The initial data processing is covered by Section 2.2.3.

on the native ICON grid. Deterministic analysis products are generated by a hybrid system named EnVar which combines variational and ensemble methods. In addition, ensemble analysis products are available from January 2018 onwards. See Chapter 11 for more information on DWD’s data assimilation system.

Choosing the Initial Data

Basically, what can be thought of as the analysis consists of two different data products: a first guess file and an analysis file. The term first guess denotes a short-range forecast of the NWP model at hand, whereas the term analysis denotes all those fields which have been updated by the assimilation system.

Several combinations of these files exist, with specific pros and cons:

Uninitialized analysis for IAU

This product consists of a first guess file and an analysis file, with the latter containing analysis increments. It is meant for starting the model in incremental analysis update (IAU) mode. IAU is a model internal filtering method for reducing spurious noise introduced by the analysis (see below).
While this initialization method performs best in terms of noise reduction, it bears
the disadvantage that the corresponding analysis product cannot be interpolated
horizontally in a straightforward manner. This prevents its use on custom target
grids. The underlying reason is that the analysis product contains tiled surface data.
Remapping of tiled data sets makes no sense, since the tile-characteristics can differ
significantly between a source and target grid cell. Only \textit{aggregated} surface fields can
be remapped (see Section 3.8.10 for more details on the surface tile approach).

\textbf{Uninitialized analysis}
This product consists of a first guess file and an analysis file, with the latter con-
taining \textit{full} analysis fields instead of increments.

The model state is abruptly pulled towards the analyzed state right before the first
time integration step. Thus, no noise filtering procedure is included. This conceptu-
ally easy approach comes at the price of a massively increased noise level at model
start. Due to the lack of tiled surface data, this product can be interpolated hori-
izontally to arbitrary custom target grids without any hassle.

\textbf{Initialized analysis}
This product consists of a single file only, containing the analyzed state. First guess
and analysis fields have already been merged and filtered by means of an asymmet-
ric IAU. The noise level introduced by this product is very moderate. In addition,
this product can also be interpolated horizontally to arbitrary custom target grids.

The level of spurious noise that emerges from each of these analysis products is compared
in Figure 2.6. It shows the area averaged absolute surface pressure tendency as a func-
tion of simulation time, which is a measure of spurious gravity-noise induced by spurious
imbalance in the initial conditions. It is defined as

\[
\left\langle \left| \frac{dp_s}{dt} \right| \right\rangle = \frac{1}{A} \sum_i \left| \frac{dp_s}{dt} \right| \Delta a_i = \frac{1}{A} \sum_i \left( \sum_k \left| -g \nabla h \cdot (\rho \hat{v}_h) \Delta z_k \right| \right) \Delta a_i,
\]

with $A$ denoting the earth’s surface and $\Delta a_i$ denoting the area of the $i$th cell. The math-
ematical steps to obtain the pressure tendency equation are discussed at the end of Sec-
tion 3.3. The blue, green and red curves show results for the \textit{uninitialized analysis for IAU},
the \textit{initialized analysis}, and the \textit{uninitialized analysis}, respectively. It can be seen that for
the \textit{uninitialized analysis} the noise level at simulation start is significantly increased when
compared to the other two modes. It takes about two days of model forecast for the noise
levels to align. While the \textit{uninitialized analysis for IAU} performs best in terms of noise-
level, it has the disadvantage that some data fields cannot easily be interpolated in the
horizontal, such that the application of this mode is typically restricted to the horizontal
grid (13 km grid spacing) used operationally at DWD.

\textbf{Important note:} For external users we strongly recommend to use the \textit{initia-
ized analysis} for model initialization, since it constitutes a good compromise
between accuracy and practicability.
Figure 2.6.: Area averaged absolute surface pressure tendency in hPa as a function of simulation time. Curves differ in terms of the way the model is initialized, with the uninitialized analysis for IAU in blue, the uninitialized analysis in red and the initialized analysis in green.

Option 1: Uninitialized Analysis Product for IAU

In the incremental analysis update (IAU) method (Bloom et al., 1996, Polavarapu et al., 2004) the analysis increment is not added in one time step completely, but it is integrated into the model integration and added to the model states $x_k^{(b)}$ over an interval $\Delta t$, which by default is $\Delta t = 3\, \text{h}$. This method of tentatively pulling the model from its current state (first guess) towards the analyzed state acts as a low pass filter in frequency domain on the analysis increments, such that small scale unbalanced modes are effectively filtered.

In the following, let us assume that we want to start a model forecast at 00 UTC. Technically, the application of this method has some potential pitfalls, which the user should be aware of:

- The analysis file has to contain analysis increments (i.e. deviations from the first guess) instead of full fields, with validity time 00 UTC. The only exceptions are \texttt{FR\_ICE} and \texttt{T\_SEA} (or alternatively \texttt{T\_SO(0)}), which must be full fields (see Table 2.1).

- The model must be started from a first guess which is shifted back in time by $1.5\, \text{h}$ w.r.t. the analysis. Thus, in the given example, the validity time of the first guess must be 22:30 UTC of the previous day. This is because “dribbling” of the analysis increments is performed over the symmetric 3 h time window [00 UTC − 1.5h, 00 UTC + 1.5h]. See Section 5.1.3 for an illustration of this process.
Table 2.1 provides an overview of the fields contained in the *uninitialized analysis product for IAU* for 00 UTC. Columns 1 to 3 show DWD’s GRIB2 shortName, the unit and a short description of the fields, respectively. Columns 3 and 4 indicate, whether the field is part of the first guess file and/or analysis file. The marker ⊗ highlights analysis increments as opposed to full fields. First guess fields which are optional for starting the model with IAU are highlighted in blue, with their scope being indicated in the description. If one or more of these fields are not available, a cold-start of these fields is performed given that the parameterizations for which they are needed are activated.

As will be explained in Section 11.2, the atmospheric analysis is performed more frequently than the surface analysis. Therefore, the analysis product at times different from 00 UTC usually contains only a subset of the fields provided at 00 UTC. Consequently, Table 2.1 will look different for non-00 UTC runs in such a way that the fields

\[ \text{FR, T, TKE, VN, } \]

will not be provided in the analysis file and will instead be contained in the first guess file.

**Table 2.1**: Content of the *uninitialized analysis product for IAU*, separated into first guess (FG) and analysis (ANA). Optional fields for model initialization are marked in blue. The marker ⊗ indicates analysis increments as opposed to full fields. Analysis fields highlighted in red are only available for 00 UTC. The validity date of the first guess is shifted back by 1.5 h w.r.t. the start date.

<table>
<thead>
<tr>
<th>shortName</th>
<th>Unit</th>
<th>Description</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>FG</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ANA</td>
</tr>
<tr>
<td>DEN</td>
<td>kg m(^{-3})</td>
<td>air density</td>
<td>×</td>
</tr>
<tr>
<td>P</td>
<td>Pa</td>
<td>pressure</td>
<td>×</td>
</tr>
<tr>
<td>QC</td>
<td>kg kg(^{-1})</td>
<td>cloud liquid water mass fraction</td>
<td>×</td>
</tr>
<tr>
<td>QI</td>
<td>kg kg(^{-1})</td>
<td>cloud ice mass fraction</td>
<td>×</td>
</tr>
<tr>
<td>QR</td>
<td>kg kg(^{-1})</td>
<td>rain water mass fraction</td>
<td>×</td>
</tr>
<tr>
<td>QS</td>
<td>kg kg(^{-1})</td>
<td>snow mass fraction</td>
<td>×</td>
</tr>
<tr>
<td>QV</td>
<td>kg kg(^{-1})</td>
<td>water vapor mass fraction</td>
<td>×</td>
</tr>
<tr>
<td>T</td>
<td>K</td>
<td>air temperature</td>
<td>⊗</td>
</tr>
<tr>
<td>TKE</td>
<td>m(^2)s(^{-2})</td>
<td>turbulent kinetic energy</td>
<td>⊗</td>
</tr>
<tr>
<td>U, V</td>
<td>m s(^{-1})</td>
<td>horizontal velocity components</td>
<td>⊗</td>
</tr>
<tr>
<td>VN</td>
<td>m s(^{-1})</td>
<td>edge normal velocity component</td>
<td>×</td>
</tr>
<tr>
<td>W</td>
<td>m s(^{-1})</td>
<td>vertical velocity</td>
<td>×</td>
</tr>
<tr>
<td>ALB_SEAICE</td>
<td>%</td>
<td>sea ice albedo</td>
<td>×</td>
</tr>
<tr>
<td></td>
<td></td>
<td>scope: lprog_albsi=.TRUE. (namelist lnd,mml)</td>
<td></td>
</tr>
<tr>
<td>C_T_LK</td>
<td>1</td>
<td>shape factor w.r.t. temp. profile in the thermocline</td>
<td>×</td>
</tr>
</tbody>
</table>

*Continued on next page*
### Table 2.1.: Continued from previous page

<table>
<thead>
<tr>
<th>Variable</th>
<th>Unit</th>
<th>Description</th>
<th>Scope</th>
</tr>
</thead>
<tbody>
<tr>
<td>EVAP_LOCAL</td>
<td>kg m(^{-2})</td>
<td>evaporation of plants (integrated since &quot;nightly reset&quot;)</td>
<td>×</td>
</tr>
<tr>
<td>FRESHSNW</td>
<td>1</td>
<td>age of snow indicator</td>
<td>×</td>
</tr>
<tr>
<td>FR_ICE</td>
<td>1</td>
<td>sea/land ice fraction</td>
<td>×</td>
</tr>
<tr>
<td>H_ICE</td>
<td>m</td>
<td>sea ice depth</td>
<td>×</td>
</tr>
<tr>
<td>H_ML_LK</td>
<td>m</td>
<td>mixed-layer thickness</td>
<td>×</td>
</tr>
<tr>
<td>H_SNOW</td>
<td>m</td>
<td>snow depth</td>
<td>×</td>
</tr>
<tr>
<td>HSNOW_MAX</td>
<td>m</td>
<td>maximum snow depth reached within current snow-cover period</td>
<td>×</td>
</tr>
<tr>
<td>QV_S</td>
<td>kg kg(^{-1})</td>
<td>surface specific humidity</td>
<td>×</td>
</tr>
<tr>
<td>RHO_SNOW</td>
<td>kg m(^{-3})</td>
<td>snow density</td>
<td>×</td>
</tr>
<tr>
<td>SNOAG</td>
<td>d</td>
<td>duration of current snow-cover period</td>
<td>×</td>
</tr>
<tr>
<td>SNOWC</td>
<td>%</td>
<td>snow cover</td>
<td>×</td>
</tr>
<tr>
<td>T_BOT_LK</td>
<td>K</td>
<td>temperature at water-bottom/sediment interface</td>
<td>×</td>
</tr>
<tr>
<td>T_G</td>
<td>K</td>
<td>surface temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_ICE</td>
<td>K</td>
<td>sea ice temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_MNW_LK</td>
<td>K</td>
<td>mean temperature of the water column</td>
<td>×</td>
</tr>
<tr>
<td>T_SEA</td>
<td>K</td>
<td>sea surface temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_SNOW</td>
<td>K</td>
<td>snow temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_WML_LK</td>
<td>K</td>
<td>mixed-layer temperature</td>
<td>×</td>
</tr>
<tr>
<td>W_I</td>
<td>kg m(^{-2})</td>
<td>water content of interception layer</td>
<td>×</td>
</tr>
<tr>
<td>Z0</td>
<td>m</td>
<td>surface roughness length</td>
<td>×</td>
</tr>
<tr>
<td>T_SO</td>
<td>K</td>
<td>soil temperature</td>
<td>×</td>
</tr>
<tr>
<td>W_SO</td>
<td>kg m(^{-2})</td>
<td>soil water content (liq. + ice)</td>
<td>×</td>
</tr>
<tr>
<td>W_SO_ICE</td>
<td>kg m(^{-2})</td>
<td>soil ice content</td>
<td>×</td>
</tr>
</tbody>
</table>

### Option 2: Uninitialized Analysis Product

The *uninitialized analysis* can be used if, for some reason, the model should be started without any noise filtering procedure. The first guess and analysis file are read in and merged by the model, i.e. the model state is abruptly pulled towards the analyzed state right before the first time integration step. This conceptually easy approach comes at the price of a massively increased noise level at the beginning of the simulation.
The validity time of the first guess and analysis must match the model’s start date. Table 2.2 provides an overview of the fields contained in the uninitialized analysis product for 00 UTC. Columns 3 and 4 again indicate, whether a specific field is contained in the first guess file and/or analysis file. Fields which are optional for starting the model are highlighted in blue, with their scope being indicated in the description. If one or more of these fields are not available, a cold-start of these fields is performed given that the parameterizations for which they are needed are activated.

As already explained in the previous section, the analysis at times different from 00 UTC will only contain a subset of the fields provided at 00 UTC. Table 2.2 will differ for non-00 UTC runs in the way that the fields FR, H, T, SEA, W, SO will not be available from the analysis file but from the first guess file.

Table 2.2: Content of the uninitialized analysis product, separated into first guess (FG) and analysis (ANA). Optional fields for model initialization are marked in blue. Analysis fields highlighted in red are only available for 00 UTC.

<table>
<thead>
<tr>
<th>shortName</th>
<th>Unit</th>
<th>Description</th>
<th>Source</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEN</td>
<td>kg m$^{-3}$</td>
<td>air density</td>
<td>×</td>
<td>ANA</td>
</tr>
<tr>
<td>P</td>
<td>Pa</td>
<td>pressure</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>QC</td>
<td>kg kg$^{-1}$</td>
<td>cloud liquid water mass fraction</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>QI</td>
<td>kg kg$^{-1}$</td>
<td>cloud ice mass fraction</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>QR</td>
<td>kg kg$^{-1}$</td>
<td>rain water mass fraction</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>QS</td>
<td>kg kg$^{-1}$</td>
<td>snow mass fraction</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>QV</td>
<td>kg kg$^{-1}$</td>
<td>water vapor mass fraction</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>T</td>
<td>K</td>
<td>air temperature</td>
<td>×</td>
<td>ANA</td>
</tr>
<tr>
<td>THETA_V</td>
<td>K</td>
<td>virtual potential temperature</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>TKE</td>
<td>m$^2$s$^{-2}$</td>
<td>turbulent kinetic energy</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>U, V</td>
<td>m s$^{-1}$</td>
<td>horizontal velocity components</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>VN</td>
<td>m s$^{-1}$</td>
<td>edge normal velocity component</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>W</td>
<td>m s$^{-1}$</td>
<td>vertical velocity</td>
<td>×</td>
<td>×</td>
</tr>
<tr>
<td>ALB_SEAICE</td>
<td>%</td>
<td>sea ice albedo</td>
<td>×</td>
<td>ANA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>scope: lprog_albsi=.TRUE. (namelist lnd_nml)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C_T_LK</td>
<td>1</td>
<td>shape factor w.r.t. temp. profile in the thermocline</td>
<td>×</td>
<td></td>
</tr>
<tr>
<td>EVAP_PL</td>
<td>kg m$^{-2}$</td>
<td>evaporation of plants (integrated since ”nightly reset”)</td>
<td>×</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>scope: itype_trvg=3 (namelist lnd_nml)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>FRESHSNW</td>
<td>1</td>
<td>age of snow indicator</td>
<td>×</td>
<td>ANA</td>
</tr>
</tbody>
</table>
Table 2.2.: Continued from previous page

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
<th>Options</th>
</tr>
</thead>
<tbody>
<tr>
<td>FR_ICE</td>
<td>sea/lake ice fraction</td>
<td>×</td>
</tr>
<tr>
<td>H_ICE</td>
<td>sea ice depth</td>
<td>×</td>
</tr>
<tr>
<td>H_ML_LK</td>
<td>mixed-layer thickness</td>
<td>×</td>
</tr>
<tr>
<td>H_SNOW</td>
<td>snow depth</td>
<td>×</td>
</tr>
<tr>
<td>HSNOw_MAX</td>
<td>maximum snow depth reached within current snow-cover period scope: itype_snowevap=3 (namelist lnd_nml)</td>
<td>×</td>
</tr>
<tr>
<td>QV_S</td>
<td>surface specific humidity</td>
<td>×</td>
</tr>
<tr>
<td>RHO_SNOW</td>
<td>snow density</td>
<td>×</td>
</tr>
<tr>
<td>SNOAG</td>
<td>duration of current snow-cover period scope: itype_snowevap=3 (namelist lnd_nml)</td>
<td>×</td>
</tr>
<tr>
<td>T_BOT_LK</td>
<td>temperature at water-bottom sediment interface</td>
<td>×</td>
</tr>
<tr>
<td>T_G</td>
<td>surface temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_ICE</td>
<td>sea ice temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_MNW_LK</td>
<td>mean temperature of the water column</td>
<td>×</td>
</tr>
<tr>
<td>T_SEA</td>
<td>sea surface temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_SNOW</td>
<td>snow temperature</td>
<td>×</td>
</tr>
<tr>
<td>T_WML_LK</td>
<td>mixed-layer temperature</td>
<td>×</td>
</tr>
<tr>
<td>W_I</td>
<td>water content of interception layer</td>
<td>×</td>
</tr>
<tr>
<td>W_SNOW</td>
<td>snow water equivalent</td>
<td>×</td>
</tr>
<tr>
<td>Z0</td>
<td>surface roughness length</td>
<td>×</td>
</tr>
<tr>
<td>T_SO</td>
<td>soil temperature</td>
<td>×</td>
</tr>
<tr>
<td>W_SO</td>
<td>soil water content (liq. + ice)</td>
<td>×</td>
</tr>
<tr>
<td>W_SO_ICE</td>
<td>soil ice content</td>
<td>×</td>
</tr>
</tbody>
</table>

Option 3: Initialized Analysis Product

The initialized analysis is strongly related to the uninitialized analysis for IAU. It is a by-product of starting the model from the latter. E.g. the initialized analysis at 00 UTC is generated by starting the model from the 22:30 UTC first guess, and adding the analysis increments over an asymmetric time window of 1.5 h width until 00 UTC. Therefore the noise level of the initialized analysis product is comparable to that of the uninitialized analysis product for IAU.

For model initialization the validity time of the initialized analysis product must match the model’s start date. Table 2.3 provides an overview of the fields contained in the initialized
analysis product for 00 UTC. Fields which are optional for starting the model are highlighted in blue. If one or more of these fields are not available, a cold-start of these fields is performed given that the parameterizations for which they are needed are switched on.

Note that this product is also suitable for initializing COSMO limited area simulations. To this end it contains the atmospheric fields $T$, $P$, $U$, $V$ rather than ICON’s set of prognostic atmospheric variables, i.e. $\text{THETA}_V$, $\text{RHO}$, $\text{VN}$. For ICON, the necessary transformation is performed automatically during startup.

For recent analysis dates (say August 2018 and later) this analysis product contains both SMI and $W_{SO}$. For previous analysis dates only $W_{SO}$ is available. ICON can read any of them, however SMI is preferred and $W_{SO}$ is the fallback. Whenever this analysis product needs to be remapped and SMI is not available, it is strongly recommended to manually convert $W_{SO}$ to SMI beforehand, since interpolating $W_{SO}$ can lead to strong numerical artifacts. See Section 2.2.3 for more details.

Table 2.3.: Content of the initialized analysis product. Fields which are optional for model initialization are marked in blue.

<table>
<thead>
<tr>
<th>shortName</th>
<th>Unit</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$P$</td>
<td>Pa</td>
<td>pressure</td>
</tr>
<tr>
<td>$QC$</td>
<td>kg kg$^{-1}$</td>
<td>cloud liquid water mass fraction</td>
</tr>
<tr>
<td>$QI$</td>
<td>kg kg$^{-1}$</td>
<td>cloud ice mass fraction</td>
</tr>
<tr>
<td>$QR$</td>
<td>kg kg$^{-1}$</td>
<td>rain water mass fraction</td>
</tr>
<tr>
<td>$QS$</td>
<td>kg kg$^{-1}$</td>
<td>snow mass fraction</td>
</tr>
<tr>
<td>$QV$</td>
<td>kg kg$^{-1}$</td>
<td>water vapor mass fraction</td>
</tr>
<tr>
<td>$T$</td>
<td>K</td>
<td>air temperature</td>
</tr>
<tr>
<td>$TKE$</td>
<td>m$^2$s$^{-2}$</td>
<td>turbulent kinetic energy</td>
</tr>
<tr>
<td>$U$, $V$</td>
<td>m s$^{-1}$</td>
<td>horizontal velocity components</td>
</tr>
<tr>
<td>$W$</td>
<td>m s$^{-1}$</td>
<td>vertical velocity</td>
</tr>
<tr>
<td>ALB_SEAICE</td>
<td>%</td>
<td>sea ice albedo</td>
</tr>
<tr>
<td>C_T_LK</td>
<td>1</td>
<td>shape factor w.r.t. temp. profile in the thermocline</td>
</tr>
<tr>
<td>EVAP_PL</td>
<td>kg m$^{-2}$</td>
<td>evaporation of plants (integrated since &quot;nightly reset&quot;)</td>
</tr>
<tr>
<td>FRESHSNW</td>
<td>1</td>
<td>age of snow indicator</td>
</tr>
<tr>
<td>FR_ICE</td>
<td>1</td>
<td>sea/lake ice fraction</td>
</tr>
<tr>
<td>H_ICE</td>
<td>m</td>
<td>sea ice depth</td>
</tr>
<tr>
<td>H_ML_LK</td>
<td>m</td>
<td>mixed-layer thickness</td>
</tr>
<tr>
<td>H_SNOW</td>
<td>m</td>
<td>snow depth</td>
</tr>
</tbody>
</table>

Continued on next page
Table 2.3.: Continued from previous page

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Unit</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>H_SNOW_MAX</td>
<td>m</td>
<td>maximum snow depth reached within current snow-cover period</td>
</tr>
<tr>
<td></td>
<td></td>
<td>scope: <code>itype_snowevap=3</code> (namelist <code>lnd_nml</code>)</td>
</tr>
<tr>
<td>QV_S</td>
<td>kg kg(^{-1})</td>
<td>surface specific humidity</td>
</tr>
<tr>
<td>RHO_SNOW</td>
<td>kg m(^{-3})</td>
<td>snow density</td>
</tr>
<tr>
<td>SNOAG</td>
<td>d</td>
<td>duration of current snow-cover period</td>
</tr>
<tr>
<td></td>
<td></td>
<td>scope: <code>itype_snowevap=3</code> (namelist <code>lnd_nml</code>)</td>
</tr>
<tr>
<td>T_BOT_LK</td>
<td>K</td>
<td>temperature at water-bottom sediment interface</td>
</tr>
<tr>
<td>T_G</td>
<td>K</td>
<td>surface temperature</td>
</tr>
<tr>
<td>T_ICE</td>
<td>K</td>
<td>sea ice temperature</td>
</tr>
<tr>
<td>T_MNW_LK</td>
<td>K</td>
<td>mean temperature of the water column</td>
</tr>
<tr>
<td>T_SNOW</td>
<td>K</td>
<td>snow temperature</td>
</tr>
<tr>
<td>T_WML_LK</td>
<td>K</td>
<td>mixed-layer temperature</td>
</tr>
<tr>
<td>W_I</td>
<td>kg m(^{-2})</td>
<td>water content of interception layer</td>
</tr>
<tr>
<td>W_SNOW</td>
<td>kg m(^{-2})</td>
<td>snow water equivalent</td>
</tr>
<tr>
<td>Z0</td>
<td>m</td>
<td>surface roughness length</td>
</tr>
<tr>
<td>SMI</td>
<td>1</td>
<td>soil moisture index</td>
</tr>
<tr>
<td>T_SO</td>
<td>K</td>
<td>soil temperature</td>
</tr>
<tr>
<td>W_SO</td>
<td>kg m(^{-2})</td>
<td>soil water content (liq. + ice)</td>
</tr>
<tr>
<td>W_SO_ICE</td>
<td>kg m(^{-2})</td>
<td>soil ice content</td>
</tr>
<tr>
<td>HHL</td>
<td>m</td>
<td>vertical coordinate half level heights</td>
</tr>
</tbody>
</table>

**Downloading Initial Conditions**

ICON initial conditions are stored within DWD’s meteorological data management system SKY. Here, for better performance, meta and binary data are stored separately: The meta data are stored in a relational database, sorted by data category and time. The binary data are stored temporarily on a hard drive and subsequently moved into the DWD’s tape archive using the archiving components in SKY.

A prerequisite for data retrieval is a valid account for the database “roma”. The database can be accessed in the following ways:

- If you already have access to DWD’s Linux cluster `lce`, please contact `datenservice@dwd.de`, in order to get additional access to the database. Data retrieval will then be possible by using either SKY’s query language directly, or by using the PAMORE command-line tool (the latter will be explained below).
- An alternative way to get the data is to use the web-based PAMORE service, see `https://www.dwd.de/DE/leistungen/pamore/pamore.html`
2. Input Data

2.2 Initial Conditions

It is meant for external users which do not have direct access to DWD’s computer systems. It requires a user account, to this end please fill the registration form

Registration form for PAMORE web service

DWD’s operational analysis and forecast products for the ICON model are being stored to the SKY database since 2015-01-20. However, the set of data fields stored is subject to continuous changes and improvements. I.e. the inclusion of additional fields has become necessary with the activation of more advanced physical parameterizations. The set of data fields of the early months is likely to be incomplete with regard to the initialization procedure that is explained in this tutorial: For example, the surface tile approach (see Section 3.8.10) has been activated no earlier than December 2015.

Data retrieval with PAMORE via command-line. PAMORE (PArallel MOdel data REtrieve from Oracle databases) is a high-level tool for the retrieval of (model) data from DWD’s meteorological data management system SKY.

In order to retrieve, for example, initial data from February 1, 2019 00 UTC, on the native ICON grid the following command lines can be used for the different analysis products:

Uninitialized analysis for IAU

Global domain with 13 km grid spacing

```
pamore -d 2019020100 -lt m -iglo_startdata -iau
```

Global domain (13 km) and nested EU domain (6.5 km)

```
pamore -d 2019020100 -lt m -iglo_eu_startdata -iau
```

Note that the EU domain lacks a separate analysis for the atmosphere. If required, it must be interpolated (horizontally) from the global domain.

Uninitialized analysis

Global domain with 13 km grid spacing

```
pamore -d date -lt m -iglo_startdata
```

Global domain (13 km) and nested EU domain (6.5 km)

```
pamore -d date -lt m -iglo_eu_startdata
```

Here, `date` must be replaced by the desired date (see above).

Initialized analysis

Global domain with 13 km grid spacing

```
pamore -d date -hstart 0 -hstop 0 -lt a \ 
     -model iglo -iglo_startdata_0
```
2. Input Data

Nested EU domain (6.5 km):

```
pamore -d date -hstart 0 -hstop 0 -lt a -model ieu -iconlam_startdata_0
```

Please note that for the nested domain the optional input field TKE is not available and that Z0 and H_SNOW are only available since 2018-03-14.

A full set of command-line options can be obtained via `pamore -h`. Alternatively, they are accessible on the web via

https://webservice.dwd.de/pamore.html

**Important note:** With the additional options

- `-ires r2b06` and
- `-enum num` where `num` specifies an ensemble member (e.g. 3) or a range of ensemble members (e.g. 3 – 8),

analysis products can also be picked from the LETKF analysis ensemble consisting of 40 members with 40/20 km horizontal grid spacing. Currently this option is **not available for the initialized analysis.**

**Data retrieval with PAMORE via web** The PAMORE web service allows the user-defined selection of model fields by navigating through a sequence of HTML forms.

For the specific task of retrieving ICON initial data, however, we strongly suggest not to take this path. Instead, you should make direct use of the above PAMORE command-lines and paste them into the HTML form. By this, you can minimize the risk of missing some fields.

After submitting your database request, the data will be extracted from the database and stored on an FTP server for download. Once your request has been processed, you will receive an e-mail with information about the FTP server address and the path to your data.

### 2.2.2. Obtaining ECMWF IFS Initial Data

Model runs can also be initialized by “external” analysis files produced by the Integrated Forecast System (IFS) that has been developed and is maintained by the European Centre for Medium-Range Weather Forecasts (ECMWF).

The ICON code contains a script for the automatic request for IFS data from the MARS data base. The Meteorological Archival and Retrieval System (MARS, see https://software.ecmwf.int/wiki/display/UDOC/MARS+user+documentation) is the main repository of meteorological data at ECMWF. A full list of recommended IFS analysis fields is provided in Table 2.4.

The script for importing from MARS is located in the subdirectory
2.2 Initial Conditions

Figure 2.7.: Screenshot of the PAMORE web service. It shows the HTML form where you can place your PAMORE command line request directly.

icon/scripts/preprocessing/mars4icon_smi

**Important note:** The script `mars4icon_smi` must be executed on the ECMWF computer system!

In order to retrieve, for example, T1279 grid data with 137 levels for the July 1, 2013, the following command line is used:

```
./mars4icon_smi -r 1279 -l 1/to/137 -d 2013070100 -O -L 1 -o 20130701.grb -p 5
```

Further options are shown by typing `./mars4icon_smi -h`

Note that prior to 2013-06-25 12 UTC, only 91 instead of 137 vertical levels were used by the operational system at ECMWF. For more information, regarding changes in the ECMWF model, see

[https://www.ecmwf.int/en/forecasts/documentation-and-support/changes-ecmwf-model](https://www.ecmwf.int/en/forecasts/documentation-and-support/changes-ecmwf-model)

Note that when initializing from “external” analysis files, ICON requires the soil moisture index (SMI) and not the volumetric soil moisture content (SWV) as input. The conversion of SWV to SMI is currently performed as part of the MARS request (`mars4icon_smi`). However, this conversion is not reflected in the variable short names. The fields containing SMI are still named SWVLx, with x denoting the surface layer index. The ICON model, however, expects them to be named SMIx. Therefore, the proper output name SMIX must be specified explicitly in the namelist `input_field_nml` of `iconremap` (see the example namelist on p. 42).
Table 2.4.: Recommended IFS analysis fields on a regular lat-lon grid, as retrieved by the script mars4icon_smi. Optional fields are marked in blue.

<table>
<thead>
<tr>
<th>shortName</th>
<th>Unit</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ECMWF</td>
<td></td>
<td></td>
</tr>
<tr>
<td>U, V</td>
<td>m s(^{-1})</td>
<td>horizontal velocity components</td>
</tr>
<tr>
<td>OMEGA</td>
<td>Pa s(^{-1})</td>
<td>vertical velocity</td>
</tr>
<tr>
<td>T</td>
<td>K</td>
<td>Temperature</td>
</tr>
<tr>
<td>FI</td>
<td>m(^2) s(^{-2})</td>
<td>geopotential (at model levels)</td>
</tr>
<tr>
<td>QV</td>
<td>kg kg(^{-1})</td>
<td>specific humidity</td>
</tr>
<tr>
<td>CLWC</td>
<td>kg kg(^{-1})</td>
<td>cloud liquid water content</td>
</tr>
<tr>
<td>CIWC</td>
<td>kg kg(^{-1})</td>
<td>cloud ice content</td>
</tr>
<tr>
<td>CRWC</td>
<td>kg kg(^{-1})</td>
<td>rain water content</td>
</tr>
<tr>
<td>CSWC</td>
<td>kg kg(^{-1})</td>
<td>snow water content</td>
</tr>
<tr>
<td>SST</td>
<td>K</td>
<td>sea surface temperature</td>
</tr>
<tr>
<td>CI</td>
<td>[0,1]</td>
<td>sea-ice cover</td>
</tr>
<tr>
<td>LNSP</td>
<td>-</td>
<td>logarithm of surface pressure</td>
</tr>
<tr>
<td>Z</td>
<td>m(^2) s(^{-2})</td>
<td>surface geopotential</td>
</tr>
<tr>
<td>TSN</td>
<td>K</td>
<td>snow temperature</td>
</tr>
<tr>
<td>SD</td>
<td>m of water eqv.</td>
<td>water content of snow</td>
</tr>
<tr>
<td>RSN</td>
<td>kg m(^{-3})</td>
<td>density of snow</td>
</tr>
<tr>
<td>ASN</td>
<td>[0,1]</td>
<td>snow albedo</td>
</tr>
<tr>
<td>SKT</td>
<td>K</td>
<td>skin temperature</td>
</tr>
<tr>
<td>STL[1/2/3/4]</td>
<td>K</td>
<td>soil temperature level 1/2/3/4</td>
</tr>
<tr>
<td>SWVL[1/2/3/4]</td>
<td>m(^3) m(^{-3})</td>
<td>soil moisture index (SMI) layer 1/2/3/4</td>
</tr>
<tr>
<td>SRC</td>
<td>m of water eqv.</td>
<td>water content of interception storage</td>
</tr>
<tr>
<td>LSM</td>
<td>[0,1]</td>
<td>land/sea mask</td>
</tr>
</tbody>
</table>

2.2.3. Remapping Initial Data to Your Target Grid

Often it is desirable to run ICON at horizontal resolutions which differ from those of the initial data. One important application are high-resolution limited area runs, which start from operational ICON forecasts or analysis. In this case horizontal remapping of the initial ICON data is necessary. Note that there is no need for vertical interpolation as a separate pre-processing step. The ICON model itself will take care of interpolation of the initial data onto the model levels, assumed that the user has provided the height level field HHL and set the appropriate namelist options (see the namelist parameter init_mode).

We shortly describe the basic steps of remapping: After the successful download, the analysis data must be interpolated from a regular or triangular grid onto the ICON target grid. To this end, the iconremap utility from the DWD ICON Tools will be used in batch mode. It is important to note that very large grids should always be processed MPI-parallel in batch mode.

A typical namelist for processing initial data has the following structure:
2. Input Data

2.2 Initial Conditions

For each of the variables to be remapped, the script contains a namelist `input_field_nml` which specifies details of the interpolation methods and the output name. In this example, the 3D temperature field and the horizontal wind components $U$, $V$ are remapped from a regular grid onto a triangular ICON grid. Variables are usually accessed through their name (character string), but note that for GRIB1 input data the correct field parameter must be provided with the namelist parameter `code`.

A detailed documentation of the ICON remap command-line options and namelist parameters can be found under `dwd_icon_tools/doc/icontools_doc.pdf`, i.e. Prill (2014). If the DWD ICON tools fail and if the cause of the error does not become clear from the error message, you may increase the output verbosity by setting the command-line options `-v`, `-vv`, `-vvv` etc.

For both, DWD analysis data and ECMWF IFS data, the DWD ICON Tools contain example run scripts which performs the remapping. These files are

**DWD initial data:** dwd_icon_tools/icontools/xce_remap_inidata  
**ECMWF IFS data:** dwd_icon_tools/example/runscripts/xce_ifs2icon.run

After adjusting the necessary file and directory names the scripts can be submitted to the PBS batch system of the Cray XC 40.

Some comments are in order for particular data fields:

Note that in this example, the GRIB2 input data file also contains the specification of the input grid. Therefore the two namelist parameters `INPUT_GRID_FILENAME` and `INPUT_FILENAME_GRB` are identical!
Fields SMI1, SMI2, SMI3, SMI4. As discussed in Section 2.2.2, care must be taken to properly rename the fields SWVLx to SMIx in the case that "external" IFS analysis files are remapped. An example namelist input_field_nml is given below:

```
&input_field_nml
  inputname = "SWVL1" ! soil moisture index layer 1
  outputname = "SMIL1"
  var_in_mask = "LSM" ! field to be used for masking
  in_mask_threshold = 0.5 ! threshold for masking values of input grid
  in_mask_below = .TRUE. ! values <= mask_threshold are masked on input grid
/
```

When remapping land surface fields, it is advisable to make use of the land sea mask information of the source grid (var_in_mask="FR_LAND" in input_field_nml). By doing so, we mask out water points so that only land points contribute to the interpolation stencil, see the example above.

**Important note:** For simplicity, the script xce_remap_inidata interpolates the soil water content W_SO directly. A more accurate and advisable approach would be to interpolate the soil moisture index SMI instead (if available\(^1\)), or to convert W_SO to SMI beforehand. A short Fortran program smi_w_so.f90 which performs this conversion ships with the ICON code and can be found in the subdirectory icon/scripts/postprocessing/tools.

If SMI instead of W_SO is provided to ICON, it will automatically be converted to W_SO during setup.

**Wind fields.** For the wind fields, the standard remapping would interpolate the samples from each component U, V separately. This approach has been chosen in the example script above. However, the standard method completely decouples the components of the vector fields. It does not take into account the fact that it is a vector-field tangent to the sphere.

Therefore ICON Tools are also capable of interpolating the edge-normal wind components \(v_n\). See the ICON Tools namelist documentation regarding the interpolation \(U, V \leftrightarrow v_n\). A special namelist parameter (RBF shape parameter) must be set for this vector field interpolation with radial basis functions.

\(^1\)Starting from 2018-03-14 (2018-07-11), DWD’s operational forecast products contain the soil moisture index SMI on the EU domain (global domain) (vv=0 output, initialized analysis).
2.3. Boundary Data Preparation for ICON-LAM

When running ICON in limited area mode (LAM), lateral boundary conditions must be provided. In real case applications these are time dependent and must be updated periodically by reading input files. To this end, forecast or analysis data sets from a driving model may be used which, however, need to be interpolated horizontally to the ICON grid first.

In this section we briefly describe the process of generating these lateral boundary conditions (LBCs). Again, the basic preprocessing steps for ICON are visualized in Figure 2.8, where the additional preprocessing of boundary data constitutes the main difference to Figure 2.5.

Boundary Data Retrieval

The raw data files which are intended to be used as LBCs must contain one of the sets of variables depicted in Figure 2.9, on either a triangular ICON grid, or a regular latitude-longitude grid.

Optional fields are marked in light-gray. The fact that different sets of variables can be used, provides some flexibility in terms of the driving model. As indicated in Figure 2.9, sets I to III are typical for ICON, COSMO and IFS, respectively.

Figure 2.8.: Basic preprocessing steps for ICON-LAM (compare to Fig. 2.5). The grid generation process is described in Sections 2.1 – 2.4. The initial data processing is covered by Section 2.2.3. Finally, the script xce_limarea for extracting the boundary data is described in Section 2.3. This preprocessing step is necessary for the limited area model ICON-LAM.
### Set I (e.g. ICON)

\[
\{ U, V \} \quad \text{or} \quad \{ VN \}
\]

\[
\{ W, \ \text{THETA}_V, \ \text{DEN}, \ \text{QV}, \ \text{QC}, \ \text{QI}, \ \text{QR}, \ \text{QS}, \ \text{HHL} \}
\]

### Set II (e.g. COSMO)

\[
U, \ V, \ W, \ T, \ \text{P}, \ \text{QV}, \ \text{QC}, \ \text{QI}, \ \text{QR}, \ \text{QS}, \ \text{HHL}
\]

### Set III (e.g. IFS)

\[
U, \ V, \ \text{OMEGA}, \ T, \ \text{PS}, \ \text{QV}, \ \text{QC}, \ \text{QI}, \ \text{QR}, \ \text{QS}, \ \text{GEOSP}
\]

**Figure 2.9.** Sets of variables that may serve as lateral boundary conditions for ICON-LAM. DWD GRIB2 short names are used. Optional fields are marked in gray. PS denotes the surface pressure, or its logarithm, and GEOSP denotes the surface geopotential. Blending of the sets is not allowed. Examples of driving models are given in brackets.

---

**Important note:** The 3D field HHL field (geometric height of model half levels above mean sea level) is constant data. It needs only be contained in the raw data file whose validity date matches the envisaged model start date.

---

Similar to the retrieval of initial conditions in Section 2.2.1, it is also possible to download the lateral boundary data from the DWD database.

The following PAMORE command retrieves DWD forecast data for a time range of 36 hours with a temporal resolution of 2 hours:

```
pamore -d 2018092212 -hstart 0 -hstop 36 -hinc 2 -model ieu -ilam boundary
```

Here, the command-line arguments `-hstart`, `-hstop`, and `-model` have already been introduced for the initial data retrieval, see p. 37. The new, limited-area-specific command-line arguments are

- `-ilam_boundary`
  This parameter enables the variable set II (see Fig. 2.9) for lateral boundary conditions.

- `-hinc hours`
  This optional argument specifies the temporal resolution of the data request.

---

**ICON-LAM Preprocessing Script**

The DWD ICON Tools contain a run script `xce_limarea` which processes a whole directory of data raw files (script variable “DATADIR”), mapping the fields onto the boundary zone of a limited area grid.
2. Input Data

2.3 Boundary Data Preparation for ICON-LAM

The output files are written to the directory specified in the variable OUTDIR. The input files are read from DATADIR, therefore this directory should not contain other files and should not be identical to the output folder.

The grid file names are specified by

\[ \text{INGRID} = "\text{input_grid_file}" \quad \# \text{file name of input grid} \]
\[ \text{LOCALGRID} = "\text{grid_file.nc}" \quad \# \text{file name of limited-area (output) grid} \]

After adjusting the necessary filenames INGRID and LOCALGRID and the input directory name DATADIR in dwd_icon_tools/icontools/xce_limarea, this script performs the steps described in the following two sections. The xce_limarea script can be submitted to the PBS batch system of the Cray XC 40.

**Step 1: Extract Boundary Region from the Local Grid File**

In the first step the above ICON-LAM preprocessing script creates an auxiliary grid file which contains only the cells of the boundary zone. This step needs to be performed only once before generating the boundary data.

We use the icosub program from the collection of ICON Tools, see Section 1.3, with the following namelist:

```plaintext
&iconsub_nml
  grid_filename = "$\{\text{LOCALGRID}\}\$",
  output_type = 4,
  lwrite_grid = .TRUE.,
/
&subarea_nml
  ORDER = "$\{\text{OUTDIR}\}/\text{grid_file_lbc.nc}\$",
  grf_info_file = "$\{\text{LOCALGRID}\}\$",
  min_refin_c_ctrl = 1
  max_refin_c_ctrl = 14
/
```

Then running the icosub tool creates a grid file `grid_file_lbc.nc` for the boundary strip. The cells in this boundary zone are identified by their value in a special meta-data field, the `refin_c_ctrl` index, e.g. `refin_c_ctrl = 1, \ldots, 14`, see Figure 2.10.

The width of the extracted boundary strip is specified by the namelist parameters `min_refin_c_ctrl` and `max_refin_c_ctrl`. The maximum allowed value for `max_refin_c_ctrl` is given by `max(refin_c_ctrl) \equiv \text{bdy_indexing_depth}`, i.e. the boundary indexing depth that has been chosen when generating the limited area grid (see Section 2.1.3). The width of the extracted boundary strip must be equal or larger than the width of the lateral nudging zone in the limited-area run for which it is foreseen. A save setting would be `max_refin_c_ctrl = \text{max(refin_c_ctrl)}`, as used in this example.

Note that step 1 is not allowed given that vertical boundary nudging is used in addition to lateral boundary nudging (`nudge_type=1` in `namelist nudging_nml`). In this case, boundary data must be provided for the entire local (limited-area) grid, rather than for a boundary strip, only (see Section 6.2).
2. Input Data

Figure 2.10.: Illustration of the ICON-LAM boundary zone. The cells are identified by their `refin_c_ctrl` index, e.g. `refin_c_ctrl = 1, \ldots, 14`.

Step 2: Creating Boundary Data

Any of the data sources explained in the Sections 2.2.1 and 2.2.2 can be chosen for the extraction of boundary data. To be more precise, boundary data originating from ICON, IFS, and COSMO have successfully been used. Data sets from other global or regional models may work as well, but have not been tested yet.

We define the following namelist for the `iconremap` program from the collection of ICON Tools. This happens automatically in our ICON-LAM preprocessing script:

```plaintext
&remap_nml
  in_grid_filename  = "${INGRID}"
  in_filename      = "input_data_file"
  in_type          = 2
  out_grid_filename= "${OUTDIR}/grid_file_lbc.nc"
  out_filename     = "${OUTDIR}/data_file_lbc.nc"
  out_type         = 2
  out_filetype     = 4
/
```

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Here, the data file `input_data_file` automatically iterates over all files in `${DATADIR}`. The parameters `in_type=2` and `out_type=2` specify that both grids correspond to triangular ICON meshes (`in_grid_filename` and `out_grid_filename`). Additionally, a namelist `input_field_nml` is appended for each of the preprocessed variables.

With respect to the output filename `data_file_lbc.nc` it is a good idea to follow a consistent naming convention. See Section 6.4.1 on the corresponding namelist setup of the ICON model.

Note that the `input_data_file` must contain only a single time step when running the `iconremap` tool. The `iconremap` tool therefore must be executed repeatedly in order to process the whole list of boundary data samples (this is automatically done within the `xce_limarea` script).

After the specification of the filenames, the `xce_limarea` script automatically adds remapping parameters for all variables of the Set I in Fig. 2.9. Regarding the HHL field an additional remark is in order: Since this variable needs only be contained in the raw data file whose validity date matches the envisaged model start date (see the remark above), we set this field as optional:

```plaintext
&input_field_nml
   inputname   = "HHL"
   outputname  = "z_ifc"
   intp_method = 3
   loptional   = .TRUE.
/
```

Afterwards, the `xce_limarea` script launches the call to the `iconremap` binary.

In this context the following technical detail may considerably speed up the preprocessing: The `iconremap` tool allows to store and load interpolation weights to and from a NetCDF file. When setting the namelist parameter `ncstorage_file` (character string) in the `iconremap` namelist `remap_nml`, the remapping weights are loaded from a file with this name. If this file does not exist, the weights are created from scratch and then stored for later use. Note that for MPI-parallel runs of the `iconremap` tool multiple files are created. Re-runs require exactly the same number of processes.

### 2.4. External Parameter Files

External parameter fields describe various properties of the Earth’s surface and atmosphere, which can be assumed to be invariant during the course of a typical NWP forecast (i.e. a couple of days). Examples are the topography, the land-sea mask, the soil type and atmospheric aerosols. Most of the fields are constant in time while some are available on a monthly basis in order to represent the seasonal cycle. They are read by the model during startup. The full list of external parameter fields is given in Table 2.5.
Table 2.5.: External parameter fields which are requested by ICON during startup (in alphabetical order). Fields marked in blue are not read by ICON in operational NWP runs. They are only requested, if the respective namelist parameter is set.

<table>
<thead>
<tr>
<th>ShortName</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AER_SS12</td>
<td>Sea salt aerosol climatology (monthly mean)</td>
</tr>
<tr>
<td>AER_DUST12</td>
<td>Total soil dust aerosol climatology (monthly mean)</td>
</tr>
<tr>
<td>AER_ORG12</td>
<td>Organic aerosol climatology (monthly mean)</td>
</tr>
<tr>
<td>AER_SO412</td>
<td>Total sulfate aerosol climatology (monthly mean)</td>
</tr>
<tr>
<td>AER_BC12</td>
<td>Black carbon aerosol climatology (monthly mean)</td>
</tr>
<tr>
<td>ALB_DIF12</td>
<td>Shortwave (0.3 – 5.0 µm) albedo for diffuse radiation (monthly mean)</td>
</tr>
<tr>
<td>ALB_UV12</td>
<td>UV-visible (0.3 – 0.7 µm) albedo for diffuse radiation (monthly mean)</td>
</tr>
<tr>
<td>ALB_NI12</td>
<td>Near infrared (0.7 – 5.0 µm) albedo for diffuse radiation (monthly mean)</td>
</tr>
<tr>
<td>DEPTH_LK</td>
<td>Lake depth</td>
</tr>
<tr>
<td>EMIS_RAD</td>
<td>Surface longwave (thermal) emissivity</td>
</tr>
<tr>
<td>FOR_D</td>
<td>Fraction of deciduous forest</td>
</tr>
<tr>
<td>FOR_E</td>
<td>Fraction of evergreen forest</td>
</tr>
<tr>
<td>FR_LAKE</td>
<td>Lake fraction (fresh water)</td>
</tr>
<tr>
<td>FR_LAND</td>
<td>Land fraction (excluding lake fraction but including glacier fraction)</td>
</tr>
<tr>
<td>FR_LUC</td>
<td>Land-use class fraction</td>
</tr>
<tr>
<td>HSURF</td>
<td>Topographic height at cell centers</td>
</tr>
<tr>
<td>LAI_MX</td>
<td>Leaf area index in the vegetation phase</td>
</tr>
<tr>
<td>NDVI_MAX</td>
<td>Normalized differential vegetation index</td>
</tr>
<tr>
<td>NDVI_MRAT</td>
<td>Proportion of monthly mean NDVI to yearly maximum (monthly mean)</td>
</tr>
<tr>
<td>PLCOV_MX</td>
<td>Plant covering degree in the vegetation phase</td>
</tr>
<tr>
<td>ROOTDP</td>
<td>Root depth</td>
</tr>
<tr>
<td>RSMIN</td>
<td>Minimum stomatal resistance</td>
</tr>
<tr>
<td>SOILTYP</td>
<td>Soil type</td>
</tr>
</tbody>
</table>

Continued on next page
2.4 External Parameter Files

Table 2.5.: Continued from previous page

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSO_STDH</td>
<td>Standard deviation of sub-grid scale orographic height</td>
</tr>
<tr>
<td>SSO_THETA</td>
<td>Principal axis-angle of sub-grid scale orography</td>
</tr>
<tr>
<td>SSO_GAMMA</td>
<td>Horizontal anisotropy of sub-grid scale orography</td>
</tr>
<tr>
<td>SSO_SIGMA</td>
<td>Average slope of sub-grid scale orography</td>
</tr>
<tr>
<td>T_2M_CL</td>
<td>Climatological 2m temperature (serves as lower boundary condition for soil model)</td>
</tr>
<tr>
<td>T_2M_CLIM</td>
<td>Climatological 2m temperature (monthly mean)</td>
</tr>
<tr>
<td>TOPO_CLIM</td>
<td>Interpolated topographic height for T_2M_CLIM data</td>
</tr>
<tr>
<td>T_SEA</td>
<td>Sea surface temperature climatology (monthly mean)</td>
</tr>
<tr>
<td>Z0</td>
<td>Surface roughness length (over land), containing a contribution from subgrid-scale orography</td>
</tr>
</tbody>
</table>

2.4.1. ExtPar Software

The ExtPar software (ExtPar – External Parameters for numerical weather prediction and climate application) is able to generate external parameters for the different models GME, COSMO, HRM and ICON. Experienced users can run ExtPar on UNIX or Linux systems to transform raw data from various sources into domain-specific data files. For ICON, ExtPar will output the fields given in Table 2.5 in the NetCDF file format and GRIB2 on the native triangular grid. For a more detailed overview of ExtPar, the reader is referred to the User and Implementation Guide of ExtPar, Asensio and Messmer (2014), and, additionally Smiatek et al. (2008, 2016).

The ExtPar preprocessor is a COSMO software and not part of the ICON training course release. Still, the ExtPar tool can be accessed via the ICON grid generator web service (see Section 2.1.4). Similar as for the grid files, for fixed domain sizes and resolutions some external parameter files for the ICON model are available for download. For the NWP release these data are provided in the NetCDF file format and GRIB2.

Topography information: Please note the following remark:

The topography contained in the ExtPar data files is not identical to the topography data which is eventually used by the model. This is because at start-up, after reading the ExtPar data, the topography field is optionally filtered by a smoothing operator. Therefore, for post-processing purposes it is necessary to specify and use the topography height topography_c (GRIB2)
2. Input Data

short name HSURF) from the model output (cf. Section 7.1 and Appendix B). The same applies to the fields DEPTH_LK, FR_LAND, FR_LAKE, and Z0.

2.4.2. Additional Information for Surface Tiles

ExtPar data files are available with and without additional information for surface tiles. See Section 3.8.10 for details on the tile approach.

ExtPar files suitable for the tile approach are indicated by the suffix _tiles. They are also applicable when running the model without tiles. ExtPar files without the suffix "_tiles", however, must only be used when running the model without tiles (ntiles = 1, namelist lnd_nml).

The data files do not differ in the number of fields, but rather in the way some fields are defined near coastal regions. Without the _tiles suffix, various surface parameters (e.g. SOILTYP, NDVI_MAX) are only defined at so-called dominant land points, i.e. at grid elements where the land fraction exceeds 50%. With the _tiles suffix, however, these parameters are additionally defined at cells where the land fraction is below 50%. By this, we allow for mixed water-land points. The same holds for the lake depth (DEPTH_LK) which is required by the lake parameterization. For files without the _tiles suffix, DEPTH_LK is only defined at dominant lake points.

2.4.3. Parameter Files for Radiation

In addition to the ExtPar products, input fields for radiation are loaded into the ICON model. These constants fields are distributed together with the model code in the subdirectory icon/data.

rrtmg_lw.nc
parameters for radiative transfer calculation used for the underlying RRTMG algorithm, thermal radiation.

ECHAM6_CldOptProps.nc
Cloud optical properties for liquid clouds at 30 wavelengths used for the underlying RRTMG algorithm.

On default, ICON expects the RRTMG parameter files to be named as above. Renaming is possible, however the modified name must then be passed into ICON via the namelist parameters lrtm_filename and cldopt_filename in the namelist nwp_phy_nml.
This chapter is devoted to a summary of ICON’s model structure. The principal components are illustrated in Fig. 3.1:

**Dynamics**

The centerpiece of the numerical weather prediction system is the *dynamical core*, which integrates the discrete equations for fluid motion forward in time. ICON’s dycore will be shortly described in Sections 3.1–3.5.

**Tracer Advection**

The dynamical core is followed by the numerical advection scheme, e.g. for humidity and cloud water. Section 3.6 focuses on the different methods available in ICON.

**Physics**

The former components are then coupled to parameterizations for processes such as convection that occur on scales too small to be resolved directly. We present a comprehensive overview of the physics parameterizations (NWP-mode) in Sections 3.7–3.8.

Finally, the chapter is concluded with the discussion of variable resolution modeling.

![Figure 3.1: ICON’s model structure. This flow chart will be revisited in detail in Fig. 3.7.](image)

Before I came here I was confused about this subject. Having listened to your lecture I am still confused. But on a higher level.

---

Enrico Fermi
3.1. Governing Equations

The equation system of the ICON model is based upon the prognostic variables suggested by Gassmann and Herzog (2008). It describes a two-component system consisting of dry air and water, where water is allowed to occur in all three phases, including precipitating drops and ice particles.

As described in Wacker and Herbert (2003), an equation set for the mixture can be derived by first introducing a reference velocity into the governing equations. The equations for momentum, mass and energy of the mixture, as given below, are then formed as a sum of the constituent-specific equation sets. The specific form of the governing equations for the mixture depends on the choice of the reference velocity.

Here we have chosen the barycentric velocity as reference velocity. It is defined as

$$ v_b = \frac{\sum_k \rho_k v_k}{\sum_k \rho_k}, $$

with the partial density $\rho_k$ of constituent $k$ and its advective velocity $v_k$. For simplicity, $v_b$ will be denoted as $v$ in the following.

In order to separate turbulent fluctuations from the mean flow, a density weighted averaging (known as Hesselberg averaging) is applied. Every field $\phi$ is decomposed into a density-weighted mean and a deviation (Hesselberg, 1925)

$$ \phi = \bar{\phi} + \phi'' $$

with

$$ \bar{\phi} = \frac{\rho \phi}{\bar{\rho}} $$

and subsequent averaging, $\bar{\phi}$ denotes the classical Reynolds average. More details on density-weighted average calculus can be found e.g. in Zdunkowski and Bott (2003).

The basic Hesselberg-averaged equation system, including the shallow atmosphere approximations, reads as follows

\[
\frac{\partial \tilde{v}_n}{\partial t} + (\tilde{\zeta} + f)\tilde{v}_t + \tilde{w} \frac{\partial \tilde{v}_n}{\partial z} = -c_{pd} \tilde{\theta}_v \frac{\partial \pi}{\partial n} - F(v_n) \tag{3.1}
\]

\[
\frac{\partial \tilde{w}}{\partial t} + \tilde{v}_h \cdot \nabla \tilde{w} = -c_{pd} \tilde{\theta}_v \frac{\partial \pi}{\partial z} - g \tag{3.2}
\]

\[
\frac{c_{pd} c_{pd}}{R_d} \frac{\partial \bar{\pi}}{\partial t} + \frac{\partial \tilde{\theta}_v}{\partial t} = c_{pd} \pi \frac{\partial \tilde{\theta}_v}{\partial t} = -c_{pd} \pi \nabla \cdot (\bar{\rho} \tilde{\theta}_v) + \overline{Q} \tag{3.3}
\]

\[
\frac{\partial \bar{\pi}}{\partial t} + \nabla \cdot (\bar{\rho} \bar{\theta}_v) = \sum_k \bar{\pi}^c_{k,\text{conv}} \tag{3.4}
\]

\[
\frac{\partial \bar{\rho} \tilde{q}_k}{\partial t} + \nabla \cdot (\bar{\rho} \tilde{q}_k \bar{\theta}_v) = -\nabla \cdot \left( J^k_k + \frac{\rho \bar{v} \tilde{q}_k}{\rho} \right) + \sigma_k \tag{3.5}
\]
3.1 Governing Equations

Prognostic equations are solved for the horizontal velocity component normal to the triangle edges $\vec{v}_n$ (3.1), the vertical wind component $\vec{w}$ (3.2), virtual potential temperature $\theta_v$ (3.3), the total density of the air mixture $\bar{\rho}$ (3.4), with

$$\bar{\rho} = \sum_k \bar{\rho}_k,$$

and mass fractions (3.5)

$$\tilde{q}_k = \tilde{\rho}_k / \bar{\rho}.$$

The index $k \in \{d, v, c, i, r, s, g\}$ represents a specific constituent of the mixture. We use

- $k = d$ for dry air,
- $k = v$ for water vapor,
- $k = c$ for cloud water,
- $k = i$ for cloud ice,
- $k = r$ for rain,
- $k = s$ for snow, and
- $k = g$ for graupel.

Further explanation of symbols and variables is given in Table 3.1. Note that the corresponding data structures containing the physics and dynamics variables are outlined in Section 8.2.

The equation system (3.1)–(3.5) is supplemented by the lower boundary conditions

$$\vec{w}\big|_s = \frac{E_v}{\bar{\rho}\big|_s} - \sum_{k\text{prec}} S_k\big|_s,$$

and the equation of state

$$\bar{p} = R_d \bar{\rho} \bar{T} (1 + \alpha),$$

with

$$\alpha = \left( \frac{R_v}{R_d} - 1 \right) \tilde{q}_v - \sum_{k \neq v, d} \tilde{q}_k.$$

When expressed in terms of the prognostic variables, the equation of state reads

$$\pi = \left( \frac{R_d \tilde{\rho}_v \bar{T}}{\bar{\rho}_0} \right) \frac{R_d}{\bar{\rho}_0},$$

In contrast to the original formulation by Gassmann and Herzog (2008), we make use of the two-dimensional rather than the three-dimensional Lamb transformation to convert
### Table 3.1.: Explanation of symbols in the model equations

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$$\frac{\partial}{\partial n}$$</td>
<td>horizontal derivative in edge-normal direction</td>
</tr>
<tr>
<td>$$\hat{K}_k$$</td>
<td>horizontal component of the kinetic energy</td>
</tr>
<tr>
<td>$$\hat{\zeta}$$</td>
<td>vertical component of relative vorticity</td>
</tr>
<tr>
<td>$$f$$</td>
<td>Coriolis parameter</td>
</tr>
<tr>
<td>$$\pi$$</td>
<td>Exner function</td>
</tr>
<tr>
<td>$$c_{pd}, c_{vd}$$</td>
<td>specific heat capacity for dry air at constant pressure/volume</td>
</tr>
<tr>
<td>$$F(v_n)$$</td>
<td>turbulent momentum fluxes</td>
</tr>
<tr>
<td>$$g$$</td>
<td>acceleration of gravity</td>
</tr>
<tr>
<td>$$\overline{Q}$$</td>
<td>diabatic heat source</td>
</tr>
<tr>
<td>$$\overline{J}_k$$</td>
<td>vertical diffusion flux for constituent $$k$$</td>
</tr>
<tr>
<td>$$\overline{\sigma}_k$$</td>
<td>internal conversion rate for $$k$$th constituent (i.e. conversion among different phases or particle forms)</td>
</tr>
<tr>
<td>$$\overline{\sigma}^{conv}_k$$</td>
<td>internal conversion rate for $$k$$th constituent due to convection only</td>
</tr>
<tr>
<td>$$\overline{\rho q''_k v''}$$</td>
<td>turbulent flux of $$k$$th partial mass fraction</td>
</tr>
<tr>
<td>$$\overline{\mathcal{E}_v}$$</td>
<td>surface evaporation flux</td>
</tr>
<tr>
<td>$$\overline{S}_k$$</td>
<td>sedimentation flux of $$k$$th constituent</td>
</tr>
<tr>
<td>$$\hat{v}^T$$</td>
<td>terminal fall velocity of $$k$$th constituent</td>
</tr>
</tbody>
</table>

the nonlinear momentum advection term into a vector-invariant form. Vector-invariant means that no gradients of vectors appear in this equation, which avoids derivatives of the coordinate basis that would otherwise arise in an arbitrary coordinate frame from the nonlinear momentum advection term.

Note that we do not explicitly solve a prognostic equation for the density of dry air. From Equation (3.6) it becomes clear that the partial density of one constituent (here $$\overline{\rho_d}$$) can be diagnosed, given that a prognostic equation for the total density and all but one partial densities is solved. The reconstructed tangential velocity component is denoted as $$\hat{v}_t$$, and in accordance with the model code, it is assumed here that $$(\hat{v}_t, \hat{v}_n, \hat{w})$$ form a right-handed system.

From the Equations (3.4), (3.5) for total density and partial densities some important constraints can be derived which must hold in the discretized analogue in order to achieve mass conservation. First of all, the total density is defined as the sum of all partial densities, as shown by Eq. (3.6). From Eq. (3.7) it follows that

$$\sum_k \hat{\rho}_k = 1.$$
Likewise, the prognostic equation for total density (3.4) should be obtained by summing the budget equations (3.5) of all constituents. As a consequence the following constraints hold:

$$\sum_k J_k = 0, \quad \sum_k \rho q_k^r \mathbf{v}^r = 0.$$

### 3.2. The Model Reference State

Dynamics in the atmosphere are characterized by small variations of thermodynamic quantities with respect to some background state. Therefore, like many other modeling frameworks, ICON makes use of an atmospheric reference state, i.e. the thermodynamic variables are defined as the sum of a reference state and a deviation from that. The reference state is assumed to be at rest and horizontally homogeneous, constant in time, dry and hydrostatically balanced.

Any grid-scale thermodynamic variable \( \hat{\psi} \) can then be written as

$$\hat{\psi}(\lambda, \phi, z, t) = \psi_0(z) + \psi'(\lambda, \phi, z, t).$$

The suffix 0 denotes the reference state while the prime denotes the grid-scale deviation. Thus, for the prognostic thermodynamic variables one gets

$$\rho(\lambda, \phi, z, t) = \rho_0(z) + \rho'(\lambda, \phi, z, t)$$

$$\pi(\lambda, \phi, z, t) = \pi_0(z) + \pi'(\lambda, \phi, z, t)$$

$$\hat{\theta}_v(\lambda, \phi, z, t) = \theta_{v0}(z) + \theta_v'(\lambda, \phi, z, t).$$

The background state components \( \rho_0, \pi_0, \) and \( \theta_{v0} \) are related by the equation of state (3.9) and are hydrostatically balanced, i.e.

$$\pi_0 = \left( \frac{R_d \rho_0 \theta_{v0}}{p_0} \right) \frac{\theta_{v0}}{c_v} \frac{\rho_0}{c_d}$$

$$\frac{d \pi_0}{dz} = -\frac{g}{c_{pd} \theta_{v0}}$$  \hspace{1cm} (3.10)

The actual vertical reference profiles can be obtained by integration of Eq. (3.10) given that suitable boundary values are provided. The reference state in ICON is identical to the state used by the COSMO model.

In a global model like ICON, the local deviation from such a horizontally homogeneous state can be quite substantial. Therefore, no attempt is made to linearize any part of the governing equations with regard to this reference state as it is done in models based on the anelastic assumption. Instead, the main effect of introducing a reference state in a global nonhydrostatic model like ICON is the removal of horizontal base-state pressure gradient terms in the equation of motion, i.e.

$$c_{pd} \theta_v \frac{\partial \pi}{\partial n} = c_{pd} \theta_v \frac{\partial \pi'}{\partial n}.$$
This reduces the computational error in the calculation of the pressure gradient force in case of sloping coordinate surfaces. Having said that, this effect is of minor importance for ICON, as by default the horizontal pressure gradient is evaluated truly horizontal along surfaces of constant height, rather than in terrain-following coordinates, along sloping coordinate surfaces (Zängl, 2012).

Changing the discretization of the horizontal pressure gradient is possible with the namelist switch igradp_method (nonhydrostatic_nml), but not recommended. Nevertheless, the reference state is still of some use for ICON. In the standard configuration of ICON, explicit use of the reference state is made when computing the advective horizontal fluxes for \( \rho \) and \( \theta_v \).

With the base state at hand, the vertical acceleration due to the pressure gradient and gravity in Eq. (3.2) is rewritten as

\[
-c_p \partial_{\theta_v} \frac{\partial \pi}{\partial z} - g = -c_p \left( \theta_{v0} + \theta'_v \right) \frac{\partial (\pi_0 + \pi')}{\partial z} = -c_p \left( \theta_{v0} \frac{\partial \pi'}{\partial z} + \theta'_v \frac{d\pi_0}{dz} \right) - c_p \theta_{v0} \frac{d\pi_0}{dz} - g
\]

The vertical momentum equation finally reads

\[
\frac{\partial \hat{w}}{\partial t} + \hat{v} \cdot \nabla \hat{w} + \frac{\partial \hat{w}}{\partial z} = -c_p \left( \theta_{v0} \frac{\partial \pi'}{\partial z} + \theta'_v \frac{d\pi_0}{dz} \right).
\]

The perturbation fields \( \pi' \) and \( \theta'_v \) are obtained from the predicted full fields and reference fields via \( \psi' = \hat{\psi} - \psi_0 \).

### 3.3. Simplifying Assumptions in the Recent Model Version

The recent model version does not account for mass loss/gain due to precipitation/evaporation. In particular, the following assumptions are made:

The term \( \sum_k \sigma_k \text{conv}_k \) on the r.h.s. of Eq. (3.4), which describes the net mass tendency of the convection parameterization, is neglected. Thus the model neither accounts for mass loss due to convective precipitation, nor mass re-distribution due to convective motions. The mass continuity equation takes the form

\[
\frac{\partial \overline{\rho \hat{w}}}{\partial t} + \nabla \cdot (\overline{\rho \hat{w}}) = 0. \tag{3.11}
\]

At the surface, the boundary condition for \( \hat{w} \), Eq. (3.8), is approximated as

\[
\overline{\rho \hat{w}}|_s = \sum_k \overline{\rho_k \hat{w}_k}|_s = 0, \tag{3.12}
\]

which in terrain-following coordinates translates to \( \overline{\rho \hat{w}}|_s = \overline{\rho \hat{w}} \cdot \nabla h \). \( h \) denotes the topography height. Thus, the simulated atmospheric system is assumed to be closed w.r.t.
3.4 Vertical Coordinates

total mass. The sedimentation fluxes at the surface $S_k|_s$, as well as the surface evaporation flux $E_v|_s$, which appear in the partial mass continuity equations, are retained. In order for Eq. (3.12) to hold, the (implicit) assumption is that the corresponding mass loss/gain due to sedimentation/evaporation is compensated by a fictitious flux of dry air across the surface.

Away from the surface, the diffusion fluxes of all airborne constituents (except for dry air) are neglected. The diffusive fluxes of all precipitating constituents, however, are taken into account. The continuity equation for the total mass is used in the form (3.11). Since Eq. (3.11) only holds if the constraint $\sum_k J_k z = 0$ holds, again, the (implicit) assumption made is that a fictitious diffusion flux of dry air counteracts the sedimentation fluxes such that the total mass in a volume moving with the barycentric velocity is conserved.

In summary, the simplifying assumptions can be characterized by the following (equivalent) statements:

- The current model version conserves the total air mass instead of the dry air mass.
- The precipitation mass sink and the evaporation mass source are neglected in the total mass budget of the model.
- The net mass gain or loss due to precipitation/evaporation does not affect the surface pressure.

The latter point becomes obvious, when writing down the (hydrostatic) pressure tendency equation and applying the approximations (3.11)–(3.12) (see also Wacker and Herbert (2003)). Starting from the hydrostatic equation $\frac{\partial p}{\partial z} = -\rho g$ it follows:

$$\frac{\partial}{\partial t} \int_{z_s}^{0} \rho \, dz = -g \int_{z_s}^{\infty} \frac{\partial \rho}{\partial t} \, dz$$

$$\frac{\partial \rho_s}{\partial t} = g \int_{z_s}^{\infty} \frac{\partial \rho}{\partial t} \, dz$$

$$\frac{\partial \rho_s}{\partial t} = -g \int_{z_s}^{\infty} \nabla \cdot (\rho \hat{v}) - \sum_k \sigma^{conv}_k \, dz$$

$$\frac{\partial \rho_k}{\partial t} = -g \int_{z_s}^{\infty} \nabla_k \cdot (\rho \hat{v}_k) \, dz + g (\rho w)|_s + g \tilde{F}^{conv}|_s$$

$$\frac{\partial \rho_s}{\partial t} = -g \int_{z_s}^{\infty} \nabla \cdot (\rho \hat{v}_h) \, dz + g (\rho w)|_s - \sum_{k \text{prec}} \sum_{k \text{prec}} \sigma^{conv}_k |_s \left( E_v - \sum_{k \text{prec}} S_k |_s \right) + g \tilde{F}^{conv}|_s,$$

with the convective surface precipitation flux

$$\tilde{F}^{conv}|_s = \int_{z_s}^{\infty} \sum_k \sigma^{conv}_k \, dz.$$

Thus, dynamical effects of the evaporation/precipitation mass source/sink are neglected. I.e. related pressure changes are ignored.
3. Model Description

Figure 3.2: Illustration of ICON’s vertical levels. With \( \text{num}_\text{lev} \) layers, there are \( \text{num}_\text{lev} + 1 \) so-called half levels. The half levels \( k - \frac{1}{2}, k + \frac{1}{2} \) enclose layer \( k \) at the centers of which are the corresponding full levels \( k \), for \( k = 1, \ldots, \text{num}_\text{lev} \). Layer 1 is at the top of the atmosphere and layer \( n \) at the bottom of the atmosphere. Half level \( \text{num}_\text{lev} + 1 \) coincides with the Earth’s surface.

3.4. Vertical Coordinates

In a nonhydrostatic model, a vertical coordinate in terms of pressure does not make sense since it cannot be taken for granted that the pressure is monotonously decreasing with increasing altitude. In general, the air mass of an air column above a certain point can not be calculated from the pressure at that point anymore. Instead, a geometric altitude grid has to be used.

In ICON the choice is a height based coordinate system that follows the terrain and consequently, the top and bottom triangle faces are inclined with respect to the tangent plane on a sphere. Due to the fact that the model levels gradually change into levels of constant height as the distance from the lower boundary increases, top and bottom triangle faces of a grid box are also slightly inclined to each other. The exact altitude of each grid box depends on the geographical position on the globe. The top and bottom faces are called half levels of the vertical grid, the center of the box is said to be at the full level of the vertical grid, see Fig. 3.2 for an illustration. Note that the numbering of full and half levels is top-down, starting with \( l = 1 \) for the top half- and full level. A Lorenz-type staggering is used in the vertical, which means that horizontal velocity, virtual potential temperature and density are defined at full levels, whereas vertical velocity is defined at half levels.
When setting up an ICON simulation, the total number of vertical levels has to be specified for each domain via

\texttt{num\_lev (namelist run\_nml, list of integer value)}

Comma-separated list of integer values giving the number of vertical full levels for each domain.

If the number of vertical levels is desired to vary between domains, setting the namelist parameter \texttt{lvert\_nest (run\_nml)} to .TRUE. is required. See Section 3.9.3 for more information on vertical nesting.

Two variants of a height-based terrain-following vertical coordinate are available in ICON. Both of which are briefly described below.

### 3.4.1. Terrain-following Hybrid Gal-Chen Coordinate

The terrain-following hybrid Gal-Chen coordinate (Simmons and Burridge, 1981) is an extension of the classic terrain-following coordinate introduced by Gal-Chen and Somerville (1975). As shown by Klemp (2011), it can be expressed in the form

\[
\begin{align*}
z(x, y, \eta) &= \frac{(H - B'(\eta) h(x, y))}{H} \eta + B'(\eta) h(x, y) \\
&= \eta + B'(\eta) \left(1 - \frac{\eta}{H}\right) h(x, y),
\end{align*}
\]

where \(z\) represents the height of the coordinate surfaces \(\eta\), \(h(x, y)\) is the terrain height, and \(H\) denotes the domain height. With \(B'(\eta) = 1\) the coordinate reverts to the classic formulation by Gal-Chen and Somerville (1975), i.e. the coordinate is terrain-following at the surface (\(\eta = 0\)) and becomes flat at model top (\(\eta = H\)). By choosing \(B'\) appropriately, a more rapid transition from terrain-following at the surface toward constant height can be achieved. One popular choice is to set

\[
B'(\eta) \left(1 - \frac{\eta}{H}\right) = 1 - \frac{\eta}{z_{\text{flat}}}, \quad \text{with } z_{\text{flat}} < H
\]

such that coordinate surfaces become constant height surfaces above \(z = z_{\text{flat}}\). Sometimes, Equation (3.13) is also written in the discretized form

\[
z_h(x, y, k) = A(k) + B(k) h(x, y), \quad k = 1, \ldots, \text{num\_lev} + 1
\]

where \(k\) denotes the vertical level index and \(z_h\) is the half level height.

### Configuring the Hybrid Gal-Chen Coordinate

The main switch for selecting the Gal-Chen hybrid coordinate is

\texttt{ivctype = 1 (namelist nonhydrostatic\_nml, integer value)}
In that case the user has to provide the vertical coordinate table (vct) as an input file. The table consists of the A and B values (see Equation (3.14)) from which the half level heights \( z_h(x, y, k) \) can be deduced. \( A(k)[m] \) are fixed height values, with \( A(1) \) defining the model top height \( H \) and \( A(\text{num lev} + 1) = 0 \text{ m} \). The dimensionless values \( B(k) \) control the vertical decay of the topography signal, with \( B(1) = 0 \) and \( B(\text{num lev} + 1) = 1 \). Thus, at each horizontal grid point \( z_h(x, y, 1) \) is the model top height, while \( z_h(x, y, \text{num lev} + 1) \) is the surface height.

The structure of the expected input file is depicted in Table 3.2. Example files can be found in `icon/vertical_coord_tables`. The file must obey the following naming rule: `atm_hyb_sz_[nlev]`, where `[nlev]` must be replaced by the total number of full levels. ICON expects the file to be located in the base directory from which the model is started. Note that there is no namelist parameter with which the location and name of the file can be specified. The namelist parameter `vertical_grid_filename (grid_nml)`, which one might be tempted to use for that purpose, is meant for something slightly different: Its purpose is to read in a vertical grid file (NetCDF format) that has been written to disc by a previous ICON run.

```plaintext
# File structure
# ----------------
# A and B values are stored in arrays vct_a(k) and vct_b(k).
# The files in text format are structured as follows:
#
# | k   vct_a(k) [m]  vct_b(k) | <- first line of file = header line
# | 1   A(1)       B(1)       | <- first line of A and B values
# | 2   A(2)       B(2)       |                      
# | 3   A(3)       B(3)       |                      
# | ...                     |                      
# | nlev+1 A(nlev+1) B(nlev+1)| <- last line of A and B values
# |=============================| <- lines from here on are ignored
# |Source: | by mo_hyb_params:read_hyb_params
# |<some lines of text>      |
# |Comments:                  |
# |<some lines of text>      |
# |References:                |
# |<some lines of text>      |
# -------------------------------------
```

Table 3.2.: Structure of vertical coordinate table as expected by the ICON model.

### 3.4.2. SLEVE Coordinate

In the case of a terrain-following hybrid Gal-Chen coordinate the influence of terrain on the coordinate surfaces decays only linearly with height. The basic idea of the *Smooth Level Vertical* SLEVE coordinate (Schär et al., 2002, Leuenberger et al., 2010) is to increase
the decay rate, by allowing smaller-scale terrain features to be removed more rapidly with height. To this end, the topography \( h(x, y) \) is divided into two components

\[ h(x, y) = h_1(x, y) + h_2(x, y), \]

where \( h_1(x, y) \) denotes a smoothed representation of \( h(x, y) \), and \( h_2(x, y) = h(x, y) - h_1(x, y) \) contains the smaller-scale contributions. The coordinate is then defined as

\[ z(x, y, \eta) = \eta + B_1(\eta) h_1(x, y) + B_2(\eta) h_2(x, y). \]

Different decay functions \( B_1 \) and \( B_2 \) are now chosen for the decay of the large- and small-scale terrain features, respectively. These functions are selected such that the influence of small-scale terrain features on the coordinate surfaces decays much faster with height than their large-scale (well-resolved) counterparts. The squeezing of the model layers above steep mountains is limited automatically in order to prevent (nearly) intersecting layers that would cause numerical instabilities.

**Configuring the SLEVE Coordinate**

The main switch for selecting the SLEVE vertical coordinate is

\[ \text{ivctype} = 2 \text{ (namelist nonhydrostatic.nml, integer value)} \]

This is the default and recommended setting. The vertical grid is constructed during the initialization phase of ICON, based on additional parameters defined in `sleve.nml`. Here we will only discuss the most relevant parameters. For a full list, the reader is referred to the namelist documentation.

**Namelist sleve.nml:**

- **top_height** *(namelist sleve.nml, real value)*
  Height of model top.

- **flat_height** *(namelist sleve.nml, real value)*
  Height above which the coordinate surfaces become constant height surfaces.

- **min_lay_thckn** *(namelist sleve.nml, real value)*
  Layer thickness of lowermost layer.

**Note for advanced users:** On default, a vertical stretching is applied such that coordinate surfaces become non-equally distributed along the vertical, starting with a minimum thickness of \( \text{min_lay_thckn} \) between the lowermost and second lowermost half-level. If constant layer thicknesses are desired, \( \text{min_lay_thckn} \) must be set to a value \( \leq 0 \). The layer thickness is then determined as \( \text{top_height}/\text{num_lev} \). Control output of the vertical layer distribution is written to `stderr`.
3.5. Temporal Discretization

In this section we will focus on time differencing in isolation and will neglect any complexity due to space differencing. Before we dive into the (nasty) details of ICON’s time discretization, let us take a step back and try to grasp the basic idea.

3.5.1. Basic Idea

Consider an arbitrary first-order ordinary differential equation of the form

$$\frac{d\psi(\vec{x}, t)}{dt} = F(\psi(\vec{x}, t), t).$$

(3.15)

Here, $\psi : \mathbb{R}^3 \times \mathbb{R} \rightarrow \mathbb{R}^q$ is the $q$-dimensional vector of state variables. In real applications, the vector-valued flux function $F : \mathbb{R}^q \times \mathbb{R} \rightarrow \mathbb{R}^q$ might be very complex.

Let $t_{n-m}$ denote some time in the past and $t_{n+1}$ some time in the future. Now we integrate Eq. (3.15) from time $t_{n-m}$ to $t_{n+1}$, which gives

$$\psi(\vec{x}, t_{n+1}) - \psi(\vec{x}, t_{n-m}) = \int_{t_{n-m}}^{t_{n+1}} F(\psi(\vec{x}, t), t) \, dt.$$

We can try to approximate the integral on the right hand side (r.h.s.) using some weighted average of $F$ at known discrete time levels. In the following we will restrict ourselves to simple two-level time-stepping schemes, i.e. we set $m = 0$ and only make use of $F$ at the discrete times $t_n$ and $t_{n+1}$ (for a general survey of time-differencing schemes, see Randall (2017)). With this restriction we get

$$\psi(\vec{x}, t_{n+1}) - \psi(\vec{x}, t_n) = \Delta t \left( \alpha F_{n+1} + \beta F_n \right).$$

(3.16)

The coefficients $\alpha$ and $\beta$ must satisfy the so called consistency condition

$$\alpha + \beta = 1,$$

such that the r.h.s. of Eq. (3.16) represents some averaged $F$. Equation (3.16) represents a whole family of simple two-step schemes. For example, by choosing $\alpha = 0$, $\beta = 1$ we arrive at the simple Euler forward scheme, while for $\alpha = 1$, $\beta = 0$ we get the (implicit) Euler backward scheme, both of which are first order accurate. Choosing $\alpha = \beta = 0.5$ leads to the implicit trapezoidal scheme, which is of second order accuracy.

One way to avoid the implicity of the trapezoidal scheme while retaining higher order is to switch to iterative schemes, also known as predictor-corrector schemes. This is the way pursued in ICON. The key idea of the predictor-corrector scheme is to replace the unwieldy $F_{n+1}$ by an estimate $F^*_n + 1 = F(\psi^*_n, t_{n+1})$ with $\psi^*_n$ computed by an explicit scheme, e.g. a forward Euler scheme. The full predictor-corrector scheme reads

predictor: \hspace{1cm} \psi^*(\vec{x}, t_{n+1}) = \psi(\vec{x}, t_n) + \Delta t F_n

corrector: \hspace{1cm} \psi(\vec{x}, t_{n+1}) = \psi(\vec{x}, t_n) + \Delta t \left\{ F^*_n + 1, F_n \right\} \alpha

(3.17)
Here we have introduced the notation
\[
\{x, y\}_\alpha := \alpha x + (1 - \alpha) y.
\]

Note that for \(\alpha = 1\), Equation (3.17) is an imitation of the Euler backward scheme (termed Matsuno scheme, (Matsuno, 1966)), while for \(\alpha = 0.5\), it is an imitation of the trapezoidal scheme (termed Heun’s method). The Matsuno scheme has first-order accuracy, and Heun’s method has second-order accuracy.

In simplified terms, the time integration scheme of ICON can be regarded as a mixture of the Matsuno scheme and the Heun scheme, as the coefficient \(\alpha\) used by ICON varies between these two extremes.

### 3.5.2. Implementation Details

Now, in the terminology of the previous section, in ICON we have
\[
\psi(\vec{x}, t) = [v_n(\vec{x}, t), w(\vec{x}, t), \rho(\vec{x}, t), \pi(\vec{x}, t)]^T.
\]

Here, we have omitted the partial densities \(\rho q_k\) from the vector of state variables. The prognostic equation (3.5) is treated with a different time discretization, as will be explained in Section 3.6.

For the sake of brevity we omit the notation for Reynolds- and Hesselberg averages. The subscripts \(n, n + 1^*\) and \(n + 1\) will be used to denote the current time level, the resulting time level of the predictor step and the new time level, respectively. They should not be confused with the subscript \(n\) used to denote the normal velocity component \(v_n\), or the horizontal derivative in edge-normal direction \(\partial/\partial n\).

The time discretization complies with the explicit two-time level predictor-corrector scheme which was described in the previous section, except for those terms describing vertical sound-wave propagation. These terms, i.e. vertical derivatives of \(w\) and \(\pi\) are treated implicitly for reasons of numerical stability and efficiency. Below, the implicit terms are marked in blue.

**Predictor step:**
\[
\begin{align*}
\frac{v_n^{n+1^*} - v_n^n}{\Delta t} &= -\text{adv}(v_n^n) - c_{pd}\theta^n_v \frac{\partial \pi^{n,n}}{\partial n} + F(v_n^n) \\
\frac{w_n^{n+1^*} - w_n^n}{\Delta t} &= -\text{adv}(w^n) - c_{pd}\theta^n_v \frac{\partial \sigma_0}{\partial z} - c_{pd}\theta^n_v \left\{ \frac{\partial \pi^{n,n}}{\partial z}, \frac{\partial \pi^{n,n}}{\partial z} \right\}_\eta \\
\frac{\rho_n^{n+1^*} - \rho_n^n}{\Delta t} &= -\nabla h \cdot \left( v_n^{n+1^*} \rho^n \right) - \left[ \frac{\partial}{\partial z} \left\{ w_n^{n+1^*}, w_n^n \right\}_\eta \rho^n \right] \\
\frac{\pi_n^{n+1^*} - \pi_n^n}{\Delta t} &= -\frac{R_d}{c_{vd}} \left( \frac{\pi^n}{\rho^n \theta^n_v} \right) \left[ \nabla h \cdot \left( v_n^{n+1^*} \rho^n \theta^n_v \right) \right. \\
&\quad \left. + \left[ \frac{\partial}{\partial z} \left\{ w_n^{n+1^*}, w^n \right\}_\eta \rho^n \theta^n_v \right] \right] + Q^n
\end{align*}
\]
3. Model Description

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with

\[
\text{adv}(v^n_n) = \frac{\partial K_h^n}{\partial n} + (\zeta^n + f) v^n_t + w^n \frac{\partial v^n_n}{\partial z}
\]

\[
\text{adv}(w^n) = v^n_h \cdot \nabla w^n + w^n \frac{\partial w^n}{\partial z}.
\]

The terms \( F(v^n_n) \) and \( Q^n \) denote diabatic momentum and exner pressure tendencies due to slow physics processes, i.e. parameterized convection, orographic and non-orographic gravity waves and radiation. See Section 3.7.1 for more details on the distinction between fast and slow physics processes.

The implicitness parameter \( \eta \) (with \( 0 \leq \eta \leq 1 \)) for the vertically implicit sound wave solver has a default value of \( \eta = 0.65 \) which is usually sufficient to ensure numerical stability in real-case applications. If required, this value can be modified with the namelist parameter \( \text{vwind} \_\text{offctr} \) (\text{nonhydrostatic} \_\text{nml}). Note that \( \text{vwind} \_\text{offctr} \) presents the off-centering \( \text{vwind} \_\text{offctr} = \eta - 0.5 \) (i.e., the default is \( \text{vwind} \_\text{offctr} = 0.15 \)). Its permissible range is given by

\[
0 \leq \text{vwind} \_\text{offctr} \leq 0.5.
\]

**Corrector step:**

\[
\frac{v^{n+1}_n - v^n_n}{\Delta t} = - \left\{ \text{adv}(v^{n+1}_n), \text{adv}(v^n_n) \right\}_\alpha
- c_{pd} \left\{ \theta^{n+1}_v, \theta^n_v \right\}_\delta \left\{ \frac{\partial \pi^{n+1}_n}{\partial z}, \frac{\partial \pi^n}{\partial z} \right\}_\delta
- F_d(v^{n+1}_n) + F(v^n_n)
\]

\[
\frac{w^{n+1}_n - w^n_n}{\Delta t} = - \left\{ \text{adv}(w^{n+1}_n), \text{adv}(w^n_n) \right\}_\alpha
- c_{pd} \left\{ \theta^{n+1}_v, \theta^n_v \right\}_\delta \left\{ \frac{\partial \pi^{n+1}_n}{\partial z}, \frac{\partial \pi^n}{\partial z} \right\}_\delta
\]

\[
\frac{\rho^{n+1}_n - \rho^n_n}{\Delta t} = - \nabla_h \cdot (v^n_n \rho^n_n) - \frac{\partial}{\partial z} \left\{ \frac{\partial \pi^{n+1}_n}{\partial z}, \frac{\partial \pi^n}{\partial z} \right\}_\delta
\]

\[
\frac{\pi^{n+1}_n - \pi^n_n}{\Delta t} = - \frac{R_d}{c_{pd} (\rho^n \theta^n)} \left[ \nabla_h \cdot (v^n_n \rho^n \theta^n) \right]
+ \frac{\partial}{\partial z} \left\{ \frac{\partial \pi^{n+1}_n}{\partial z}, \frac{\partial \pi^n}{\partial z} \right\}_\delta
+ Q^n
\]

The term \( F_d(v^{n+1}_n) \) represents 4th order divergence damping which has been introduced in order to control checkerboard noise.

The parameter \( \alpha \) denotes the Matsuno parameter which was introduced in Section 3.5.1. With respect to the velocity, it can be used to make the explicit part of the corrector step...
3. Model Description

3.5 Temporal Discretization

resemble either a Matsuno-type scheme ($\alpha \rightarrow 1$) or Heun’s method ($\alpha \rightarrow 0.5$). The default value of the Matsuno-parameter for velocity is given by $\alpha = 0.75$.

A second Matsuno parameter $0 \leq \delta \leq 1$ exists for the thermodynamic variables $\rho$ and $\theta_v$, where the default value of the coefficient is $\delta = 0.4$. The corresponding namelist parameters are named veladv_offctr and rhothera_offctr in the namelist nonhydrostatic.nml. Note again that both define the off-centering from 0.5 rather than the absolute value.

In order to close the system of equations, for both substeps (predictor and corrector), the quantity $\theta_v$ must be calculated from the state variables $\pi$ and $\rho$. We explain this in the following, using the notation $\pi^{\text{new}} = \pi^{n+1}$ or $\pi^{\text{new}} = \pi^{n+1}$ for the predictor and corrector step.

Once the updated state vector

$$\psi^{\text{new}}(\vec{x}, t) = [v_n^{\text{new}}(\vec{x}, t), w_n^{\text{new}}(\vec{x}, t), \rho_n^{\text{new}}(\vec{x}, t), \pi_n^{\text{new}}(\vec{x}, t)]^T$$

is known, the virtual potential temperature $\theta_v^{\text{new}}$ is diagnosed from the linearized equation of state (3.9)

$$\frac{\pi^{\text{new}} - \pi^n}{\pi^n} = \frac{R_d (\rho \theta_v)^{\text{new}} - (\rho \theta_v)^n}{c_{vd}}.$$

(3.22)

The rationale behind that is as follows: Multiplying (3.22) by $\Delta t^{-1}$ and rearranging yields

$$\frac{(\rho \theta_v)^{\text{new}} - (\rho \theta_v)^n}{\Delta t} = c_{vd} \left( \frac{\rho^n \theta_v^n}{\pi^n} \right) \frac{\pi^{\text{new}} - \pi^n}{\Delta t}.$$

(3.23)

Thus, diagnosing $\theta_v^{\text{new}}$ from (3.22) can be interpreted as solving two prognostic equations for the two thermodynamic variables $\pi$ and $\theta_v$, instead of one prognostic equation for $\pi$ and the equation of state. By doing so, $\rho \theta_v$ is exactly conserved (in the absence of diabatic terms). The two prognostic equations are constrained by (3.23), i.e. the same flux divergence is used. As a side effect, however, the solution at a particular point in time is not constrained by the equation of state (3.9) – only its time evolution is. This approach was first described by Gassmann (2013).

Pragmatic Simplifications

A potential disadvantage of predictor-corrector schemes as compared to non-iterative schemes is its computational expense. This is because the forcing terms (l.h.s.) must be evaluated twice per time step. Therefore, several terms have been simplified provided that the simplification proved to not degrade the quality of the results significantly. The following pragmatic simplifications have been performed:

- The horizontal and vertical momentum advection at the predictor step is re-used from the corrector step of the preceding time step. With time level $n^*$ denoting $n + 1^*$ from the preceding time step this can be written as

Equation (3.18): $\text{adv}(v_n^n) \simeq \text{adv}(v_n^{n^*})$

Equation (3.19): $\text{adv}(w_n^n) \simeq \text{adv}(w_n^{n^*})$

65
By this, in each time step the momentum advection needs to be computed only once.
The first predictor step following the physics step represents an exception. At this
time level the momentum advection from the preceding corrector step does not
provide a suitable estimate, as the physics step might have changed \( v_n \) considerably.

- Another simplification relates to the horizontal pressure gradient term which occurs
in the predictor and corrector step of \( v_n \). Using time level \( n \) in the predictor and an
interpolated value between \( n \) and \( n+1 \) in the corrector provides an effective damping
mechanism for horizontally propagating sound waves, without significantly impacting
gravity waves. A very similar effect can be achieved by using the same horizontal
pressure gradient in the predictor and corrector, however, with the pressure being
extrapolated in time.

\[
\text{Equation (3.18): } c_{pd}\theta_v \frac{\partial \pi_{t,n}}{\partial n} \simeq c_{pd}\theta_v \frac{\partial \pi_{t,n}}{\partial n}
\]

\[
\text{Equation (3.21): } c_p \left\{ \theta_v^{n+1}, \theta_v^n \right\} \delta \left\{ \frac{\partial \pi_{t,n+1}^\prime}{\partial n}, \frac{\partial \pi_{t,n+1}^\prime}{\partial n} \right\} \simeq c_{pd}\theta_v \frac{\partial \pi_{t,n}}{\partial n}
\]

By this, in each time step the horizontal pressure gradient needs to be computed only
once. The time level \( \tilde{n} \) at which the horizontal gradient is taken is an extrapolated
time level using the levels \( n \) and \( n-1 \):

\[
\pi_{t,n} = (1 + \gamma)\pi_{t,n} - \gamma\pi_{t,n-1}
\]

The temporal extrapolation factor is chosen from the range \( \gamma \in \left[ \frac{1}{3}, \frac{2}{3} \right] \), with the
default being \( \gamma = 1/3 \). The corresponding namelist parameter is named \texttt{exner_expol}
\texttt{(nonhydrostatic.nml)}.

**Vertically Implicit Solver**

The solution to the predictor-corrector scheme described above is mostly straightforward,
as the majority of terms are treated in an explicit manner. This, however does not hold for
the prognostic equation for vertical wind, since the solution for \( w_{n+1}^{n+1} \) depends on \( \pi_{n+1}^{n+1} \),
which itself depends on \( w_{n+1}^{n+1} \) (see Equations (3.19) and (3.20)).

Before the exact solution method is derived below, the overall solution strategy can be
described as follows: First, \( \pi \) is eliminated from Eq. (3.19) by inserting
the prognostic equation (3.20). Then this results in a linear system of equations for the unknown
\( w \)’s in the vertical direction. The derivation for the corrector step is basically identical and differs
only w.r.t. the time levels that are used for \( \rho \) and \( \theta_v \).

We start with the vertical discretization of (3.19) and (3.20). When using basic centered
differences, and noting that \( \partial \pi / \partial t = \partial \pi' / \partial t \) this leads to

\[
w_{k+1/2}^{n+1} = Z_{k+1/2}^{\text{expl}} - \Delta t c_{pd}\theta_v^{n} \frac{\pi_{k}^{t,n+1} - \pi_{k+1}^{t,n+1}}{\Delta z_{k+1/2}}
\]

\[
\pi_{k}^{t,n+1} = Z_{k}^{\text{expl}} - \Delta t R_d \left( \rho_{k}^{t,n} \theta_v^{n} \right) \frac{(w_{k+1/2}^{n+1} \rho_{k}^{n+1} \theta_v^{n})_{k+1/2} - (w_{k+1/2}^{n} \rho_{k}^{n} \theta_v^{n})_{k+1/2}}{\Delta z_k}
\]
with the shorthand notations $Z_{k+1/2}^{expl}$ and $Z_k^{expl}$ for the explicit parts

$$Z_{k+1/2}^{expl} = \pi_k^n - \Delta t \left[ \text{adv}(w_{n+1/2}) + c_{pd} \frac{\partial \theta_{v,k+1/2}}{\partial z} \right]_{k+1/2}$$

and

$$Z_k^{expl} = \pi_k^n - \Delta t \left[ \frac{\partial k}{\partial z} \left( \frac{\Gamma_k}{\Delta z_k} \right) \right]_{k+1/2}$$

and

$$Z_k^{expl} = \pi_k^n - \Delta t \left( \frac{\partial R_d}{\partial z} \right)_{k+1/2}$$

$$= \Delta t \left[ \Delta t R_d \left( \frac{\pi_k^n}{\rho_k^n \theta_{v,k}} \right) \right]_{k+1/2}$$

$$= \Delta t \left[ \Delta t R_d \left( \frac{\pi_k^n}{\rho_k^n \theta_{v,k}} \right) \right]_{k+1/2}$$

$$= \Delta t \left[ \Delta t R_d \left( \frac{\pi_k^n}{\rho_k^n \theta_{v,k}} \right) \right]_{k+1/2}$$

$$= \Delta t \left[ \Delta t R_d \left( \frac{\pi_k^n}{\rho_k^n \theta_{v,k}} \right) \right]_{k+1/2}$$

As an intermediate step, we compute $\pi'_k^{n+1} - \pi'_k^{n+1}$ from Eq. (3.25):

$$\pi'_k^{n+1} - \pi'_k^{n+1} = Z_k^{expl} - Z_k^{expl} - \Delta t R_d \left( \frac{\pi_k^n}{\rho_k^n \theta_{v,k}} \right)_{k+1/2} \Gamma_k^n \Delta z_k$$

$$= \left( w_{n+1} \rho_{n+1} \theta_{v,k} \right)_{k+1/2} - \left( w_{n} \rho_{n} \theta_{v,k} \right)_{k+1/2}$$

$$= \left( w_{n+1} \rho_{n+1} \theta_{v,k} \right)_{k+1/2} - \left( w_{n} \rho_{n} \theta_{v,k} \right)_{k+1/2}$$

$$= \left( w_{n+1} \rho_{n+1} \theta_{v,k} \right)_{k+1/2} - \left( w_{n} \rho_{n} \theta_{v,k} \right)_{k+1/2}$$

$$= \left( w_{n+1} \rho_{n+1} \theta_{v,k} \right)_{k+1/2} - \left( w_{n} \rho_{n} \theta_{v,k} \right)_{k+1/2}$$

$$= \left( w_{n+1} \rho_{n+1} \theta_{v,k} \right)_{k+1/2} - \left( w_{n} \rho_{n} \theta_{v,k} \right)_{k+1/2}$$

$$= \left( w_{n+1} \rho_{n+1} \theta_{v,k} \right)_{k+1/2} - \left( w_{n} \rho_{n} \theta_{v,k} \right)_{k+1/2}$$

Here we have introduced $\Gamma_k^n$ as an abbreviation for

$$\Gamma_k^n = \frac{\pi_k^n}{\rho_k^n \theta_{v,k}}.$$
Equation (3.27) defines a linear system of equations from which the unknown vertical velocities \( w_{n+1}^{k-1/2} \) can be computed. The system is tridiagonal and can be solved with the Thomas algorithm (Press et al., 2007, p. 57), given that suitably defined boundary conditions are provided. So far it is assumed that the upper and lower boundary are impermeable w.r.t. to mass, i.e. we set \( w_{n+1}^{1/2} = w_{n+1}^{nlev+1/2} = 0 \). In case of vertical nesting, \( w \) at the upper boundary is interpolated from the parent domain.

3.6. Tracer Transport

Physically speaking, the tracer transport module predicts the large-scale re-distribution of tracers (e.g. water substances, aerosols, chemical species, ...) caused by air-motions. Mathematically (numerically) this is done by solving the partial mass continuity equation (3.5) for each tracer, with sources and sinks being neglected, i.e.

\[
\frac{\partial \bar{\rho} q_k}{\partial t} + \nabla \cdot (\bar{\rho} q_k \bar{v}) = 0 \tag{3.28}
\]

The numerical solution of Eq. (3.28) is based on so-called space-time finite volume methods. By space-time methods we refer to methods where the temporal and spacial discretizations are combined rather than separated. Space-time methods are also known as cell-integrated semi-Lagrangian schemes. As will become clear, such schemes are neither purely semi-Lagrangian, nor Eulerian in the classical sense. They are Eulerian in the sense that the flux of mass through the stationary walls of grid cells is considered. They are, however, semi-Lagrangian in the sense that trajectory calculations are needed for flux computation. In the literature such schemes are sometimes termed Flux Form Semi-Lagrangian (FFSL). The specific implementation in ICON partly builds upon work by Lauritzen et al. (2010, 2011a), Harris and Lauritzen (2010), Skamarock and Menchaca (2010), Miura (2007) for the horizontal and Colella and Woodward (1984) for the vertical.

As we are dealing with a Finite Volume (FV) discretization, it is worth noting that in the following all scalar variables \( \psi \), whose storage location is at the triangle cell circumcenter, are interpreted as cell averages rather than point values, i.e.

\[
\bar{\psi}_i^n = \frac{1}{\Delta V_i} \iiint_{V_i} \psi(x, y, z, t^n) \, dV,
\]

with \( \Delta V_i \) denoting the volume of the \( i^{th} \) prismatic cell. Here and in the reminder of this Section, the overbar denotes volume averages rather than Reynolds averages.

3.6.1. Directional Splitting

By integrating the continuity equation (3.28) in space over a prismatic grid cell (the so-called control volume) and in time over the time step \( \Delta t \), a solution to (3.28) can formally be written as

\[
\bar{\psi}_{i,k}^{n+1} = \bar{\psi}_{i,k}^n + \Delta t \left[ H(q^n) + V(q^n) \right], \tag{3.29}
\]
where $\mathcal{H}$ and $\mathcal{V}$ denote the horizontal and vertical transport operators acting on $q^n$, and $\rho q_{i,k}^{n+1}$ denoting the updated cell averaged partial density of constituent $k$ at the time $t^{n+1}$.

Instead of solving this somewhat unwieldy equation in one sweep, a fractional step approach is taken in ICON such that separate equations for horizontal and vertical transport are solved consecutively. Of course, replacing equation (3.29) by some approximation involving the two subproblems

$$\rho q_{i,k}^* = \rho q_{i,k}^n + \Delta t \mathcal{V}(q^n)$$
$$\rho q_{i,k}^{n+1} = \rho q_{i,k}^* + \Delta t \mathcal{H}(q^*)$$

will inevitably result in a residual error. This error is known as the splitting error. On default, the following approximation to (3.29) is used:

$$\rho q_{i,k}^* = \rho q_{i,k}^n + \Delta t \mathcal{V}(q^n)$$
$$\rho q_{i,k}^{n+1} = \rho q_{i,k}^* + \Delta t \mathcal{H}(q^*)$$

In order to maintain $O(\Delta t^2)$ accuracy, the order of the operators is reversed on alternate time steps. This might be regarded as a poor man’s Strang splitting (Strang, 1968). Full Strang-splitting of the form $[\mathcal{V}(\Delta t/2)][\mathcal{H}(\Delta t)][\mathcal{V}(\Delta t/2)]$ is also available as an option by choosing $\text{istrang=.FALSE.}$. (namelist transport_nml). Except for being more expensive (the vertical operator is called twice) no significant impact on the model results has been noted so far.

A shortcoming of the splitting (3.30), (3.31) is that it does not preserve an initially uniform tracer field (e.g. $q(x,y,z,t_0) = 1$) in a deformational non-divergent flow since there is not enough information available to correctly do the conversion $\rho q^* \rightarrow \bar{q}^*$. To do so, it is necessary to keep track of the changes in partial density $\rho q$ that are solely a result of mass convergence/divergence in the directions of splitting. Therefore we follow the method of Easter (1993) wherein the mass continuity equation (3.5) is simultaneously reintegrated with that for partial mass:

$$\rho q_{i,k}^* = \rho q_{i,k}^n + \Delta t \mathcal{V}(\bar{q}^n)$$
$$\rho q_{i,k}^{n+1} = \rho q_{i,k}^* + \Delta t \mathcal{H}(\bar{q}^*)$$
$$\bar{q}_{i,k}^* = \frac{\rho q_{i,k}^*}{\rho_{i,k}^*}$$

Changes in partial density solely due to mass convergence/divergence are corrected for in equations (3.32) and (3.33). The key point here is that the intermediate density $\rho q^*$ rather than $\rho q^{n+1}$ or $\rho q^n$ is used to recover the mass fraction $\bar{q}^*$ in (3.32). For re-integration of the mass continuity equation, the mass fluxes that have already been computed in the dynamical core are re-used.
3.6.2. Horizontal Transport

A rigorous derivation of the horizontal transport operator \( H(q) \) is beyond the scope of this document. We will merely concentrate on the general concept and illustrate graphically how the scheme works. The horizontal transport scheme belongs to the class of so-called Flux Form Semi-Lagrangian (FFSL) schemes (Harris and Lauritzen, 2010). In the literature such schemes are sometimes alternatively termed Incremental remapping schemes (Lipscomb and Ringler, 2005) or streamline subgrid integration method (Yeh, 2007).

Graphical Interpretation

Figure 3.3 provides a graphical interpretation of the FFSL-scheme. Black solid lines show the triangular grid, with thick solid lines highlighting an arbitrary cell with area \( \Delta A_i \) for which the scheme will be explained. In the following we will refer to this cell as the Eulerian control volume (CV). The basic task is to compute the updated value \( \rho q_{i}^{n+1} \) for that cell on the basis of the old values \( \rho q_{i}^{n} \) and the velocity fields \( \mathbf{v}^{n} \) and \( \mathbf{v}^{n+1} \).

In order to set the stage, let us first take the Lagrangian viewpoint: Assume that the time dependent velocity field is known analytically such that we know the trajectories for all the air parcels which terminate at the walls of the Eulerian CV at the new time \( t^{n+1} \). As an example, trajectories for the air parcels terminating at the CV vertices at \( t^{n+1} \) are depicted as gray lines. Accordingly we know the position of these air parcels at time \( t^{n} \) which we will denote as the starting points. By connecting the starting points we can construct the gray shaded area known as the Lagrangian CV. The latter encompasses all air parcels that are transported into the Eulerian CV (i.e. the grid cell) during the time interval \([t^n, t^{n+1}]\). In a standard semi-Lagrangian scheme the key task is to compute an estimate of the Lagrangian CV (gray shaded), followed by a computation of the total tracer mass contained. Then, the solution \( \overline{\rho q}_{i}^{n+1} \) can easily be deduced from the Lagrangian finite-volume form of the continuity equation (3.28)

\[
\overline{\rho q}_{i}^{n+1} \Delta A_i = \overline{\rho q}_{i}^{n} \Delta a_i ,
\]

where \( \Delta A_i \) and \( \Delta a_i \) denote the area of the Eulerian and Langrangian CV, respectively (see e.g. Lauritzen et al., 2011b). \( \overline{\rho q}_{i}^{n} \) is the average tracer mass over the Lagrangian CV area \( a_i \)

\[
\overline{\rho q}_{i}^{n} = \frac{1}{\Delta a_i} \int \rho^n(x,y)q^n(x,y)dA .
\]

As mentioned before, a more Eulerian rather than semi-Lagrangian viewpoint is taken in ICON. Here we keep track of the flux of mass passing the Eulerian cell walls rather than the mass in the Lagrangian CV. This is where the yellow areas in Figure 3.3 enter the game. We will refer to these as flux areas. Since the individual edges of the Lagrangian CV pass through the flux areas during \([t^n, t^{n+1}]\), it is exactly the mass inside in the flux areas that is swept across the Eulerian CV walls during one time step. Thus, starting from the mass \( \rho q_{i}^{n} \) in the Eulerian CV and assuming that we know the shape as well as the tracer mass contained in the (yellow) flux areas, we can compute the updated value \( \overline{\rho q}_{i}^{n+1} \).
3. Model Description

3.6 Tracer Transport

Figure 3.3.: Graphical illustration of the FFSL scheme. Black solid lines show the triangular grid, with thick solid lines highlighting the Eulerian control volume under consideration. Grey area shows the Lagrangian control volume and yellow areas show the flux areas (departure region) for each cell wall.

Mathematically the scheme can be cast into the following form:

$$
\overline{\rho q}_{i}^{n+1} = \overline{\rho q}_{i}^{n} - \frac{1}{\Delta t} \sum_{e=1}^{N_e} s_{ie} F_{ie} \quad \text{with} \quad F_{ie} = \langle \overline{p_{ie}} \rangle \int_{a_{ie}} q^{n}(x,y) \, da,
$$

(3.34)

where $F_{ie}$ defines the total mass crossing the $e^{th}$ wall during $\Delta t$ and $a_{ie}$ denotes the flux area for the $e^{th}$ wall. $s_{ie} = \pm 1$ distinguishes inward and outward directed fluxes.

Note that this Eulerian viewpoint is fully equivalent to the semi-Lagrangian viewpoint. It can be shown (Lauritzen et al., 2011b) that all areas involved in our quasi-Eulerian approach (Eulerian CV plus flux areas) sum up to the Lagrangian CV.

**Basic Algorithm**

The numerical algorithm which solves Eq. (3.34) for a single Eulerian CV proceeds in 4 major stages:

1. The flux area $a_{ie}$ for each cell wall is reconstructed by means of backward trajectories.
2. For each Eulerian CV the unknown tracer subgrid distribution $q(x,y,t_0)$ is estimated from the known cell averages $\overline{q}_{i}^{n}$ of the CV itself and surrounding cells. Several polynomial reconstructions, from linear to cubic, are available.
3. The total mass $F_{ie}$ crossing the $e^{th}$ wall is estimated by numerically evaluating the integral in Eq. (3.34). I.e. the estimated subgrid distribution $q(x,y,t_0)$ is integrated over the approximated departure region $a_{ie}$ by means of Gauss quadrature.
4. The sum on the r.h.s. of Eq. (3.34) is evaluated which leads to the solution $\overline{\rho q}_{i}^{n+1}$.

3.6.3. Vertical Transport

A rigorous derivation of the vertical transport operator $V(q)$ is beyond the scope of this document. As for the horizontal operator $H(q)$ we will concentrate on the basic concept.
3. Model Description

**Figure 3.4.** The Piecewise Parabolic Method (PPM). Left: Unknown subgrid distribution \( q(z) \) gets approximated by piecewise parabolic interpolants which are \( C^0 \) continuous at cell walls. Right: Polynomial reconstruction is filtered (optional) to render the scheme monotonous. Integration step: Sub-grid distribution is integrated over the “area” \( w \Delta t \) (dark blue) in order to determine the mass which enters the \( k^{th} \) cell during \( \Delta t \).

The vertical transport scheme is based on the Piecewise Parabolic Method (PPM) developed by Colella and Woodward (1984). It is a finite volume scheme and thus inherently mass conserving. It makes use of a piecewise parabolic function for approximating the unknown subgrid distribution of a 1D scalar field \( q(z) \). The function is forced to be continuous at cell interfaces. Its construction is based on the known cell averages \( \bar{q}_k \). The PPM scheme bears some conceptual resemblance to the horizontal FFSL transport scheme. The basic concept is depicted in Figure 3.4.

**Step 1:** The subgrid distribution \( q(\zeta, k) \) is reconstructed cell-wise in a vertical column by using the parabolic interpolant

\[
q(\zeta, k) = a_0 + a_1 \zeta + a_2 \zeta^2, \quad \text{with} \quad \zeta = \frac{z - z_{k+1/2}}{\Delta z_k}.
\]  

(3.35)

\( \zeta \) is a dimensionless coordinate which is 1 at the grid cell top and 0 at its bottom. Specific to the PPM scheme is the way how this parabola is constructed. The unknown coefficients \( a_i \) are derived from the 3 constraints

\[
\int_0^1 q(\zeta, k) d\zeta = \bar{q}_k, \\
q(\zeta = 1, k) = q_u = q_{k-1/2}, \\
q(\zeta = 0, k) = q_l = q_{k+1/2},
\]

(3.36)

which, vividly said, state that the polynomial must be mass conserving, and that the polynomial equals \( q_u \) and \( q_l \) at its upper and lower end, respectively. \( q_u = q_{k-1/2} \) and \( q_l = q_{k+1/2} \) are appropriately reconstructed estimates at the upper and lower half levels and are shared between adjacent cells. By this, continuity of the reconstruction across cells is enforced.
3.6 Tracer Transport

By applying the constraints (3.36) the parabolic interpolant (3.35) can finally be written as a function of the grid scale variables \( q_k, q_{k+1/2}, q_{k-1/2} \)

\[
q(\zeta, k) = \bar{q}_k - \Delta q_k \left( \frac{1}{2} - \zeta \right) - a_{6,k} \left( \frac{1}{6} - \zeta + \zeta^2 \right),
\]

with

\[
\Delta q_k = q_{k-1/2} - q_{k+1/2}
\]

\[
a_{6,k} = 6q_k - 3q_{k-1/2} - 3q_{k+1/2}
\]

The overall accuracy of the parabolic interpolant (3.37) strongly depends on the accuracy to which the edge values \( q_{k\pm1/2} \) are known. Edge estimates must be of at least third-order in order for the interpolant to exactly reconstruct a parabola. Here we follow Colella and Woodward (1984) and compute fourth-order edge value estimates, as this turned out to be beneficial to the overall accuracy of the scheme (see Lauritzen et al. (2011b, p. 228)).

An in-depth derivation of the edge-values is beyond the scope of this tutorial. Here we simply note that the edge estimate \( q_{k+1/2} \) is computed from a cubic polynomial \( c(z) \) evaluated at \( z_{k+1/2} \). The cubic polynomial is constructed from the constraint that it must be mass conserving in each of the 4 cells surrounding half level \( z_{k+1/2} \) (likewise for \( q_{k-1/2} \), see e.g. Zerroukat et al. (2002)).

Step 2: As depicted in Figure 3.4, it is not guaranteed that the reconstruction preserves monotonicity or positive definiteness, especially near strong gradients. The scheme can optionally be made (semi-) monotonic or positive definite by filtering the polynomial reconstruction. The effect of a monotonic filter on the reconstructed parabolas is schematically depicted in Figure 3.4b. The filtering is controlled with the namelist switch \( \text{itype_vlimit (transport_nml)} \).

Step 3: In a last step, the mass that is swept across the cell wall during \( \Delta t \) is computed by integrating the subgrid distribution \( q(\zeta, k) \) over the (upwind) flux area.

\[
F_{k-1/2} = \frac{1}{\Delta t} \int_{z_{k-1/2} - w_{n+1/2} \Delta t}^{z_{k-1/2}} \rho(z)q(z)dz, \text{ for } w > 0
\]

Vividly speaking, an estimate of the flux area can be gained by launching a backward trajectory at the given cell wall. In this 1D scheme, the flux area for the cell wall at \( z_{k-1/2} \) is simply given as \( -w_{n+1/2} \Delta t \), where \( w \) is the vertical velocity provided by the dynamical core.

For \( w > 0 \) integration of (3.38) finally leads to the time-averaged vertical flux

\[
F_{k-1/2} = \rho_{k-1/2}w_{k-1/2} \left[ \bar{q}_k + \frac{1}{2} \Delta q_k (1 - C_{k-1/2}) - \frac{1}{6} a_{6,k} \left( 1 - 3C_{k-1/2} + 2C_{k-1/2}^2 \right) \right],
\]

with the Courant number \( C_{k-1/2} = w_{k-1/2} \Delta t/\Delta z_k \).

In its base version, the PPM scheme has a Courant number limitation of \( |C| \leq 1 \). It can, however, be easily extended to larger Courant numbers by breaking down the computation of the mass fluxes (3.38) into so-called integer and fractional fluxes (Lin and Rood, 1996). In its current form in ICON, the PPM scheme is stable up to \( |C| = 5 \).
3.6.4. Reduced Calling Frequency

Given that explicit time stepping is used, the continuity equation for air, the momentum and thermodynamic equation must obey the time-step restrictions imposed by the fastest waves in the system (i.e. sound waves). While the continuity equation for air is inherently coupled to the momentum equations, passive tracer transport equations can be solved in isolation given prescribed winds and air densities.

Continuity equations for passive tracers lack fast wave modes (sound and gravity waves) and, thus, have less restrictive time step limitations. Given the large number of passive tracers in state of the art climate and NWP models, significant computational cost savings can be obtained by sub-cycling the solution of the density, momentum and thermodynamic equation with respect to the tracer equations. Stated in another way, the tracer equations can be integrated with a much larger time step. In doing so, care has to be taken in order to maintain tracer-mass consistency.

In ICON, the number of times by which the integration of the continuity equation for air (and the entire dynamical core) is sub-cycled with respect to the tracer mass continuity equations is typically 5 (see Section 3.7.1). In order to preserve tracer-mass consistency, the time-averaged rather than the instantaneous mass flux is passed to the transport module. Thus, $\langle \rho_i \rangle$ in Eq. (3.34) can be expressed in terms of the time-averaged horizontal mass flux $\langle F_{xe}^m \rangle$ as

$$\rho_i = \langle F_{xe}^m \rangle \Delta t l_{ie}$$

with $\Delta t$ denoting the time step for tracer transport and $l_{ie}$ denoting the length of the $e^{th}$ cell wall.

3.6.5. Some Practical Advice

Here we give some practical guidance on how to configure the tracer transport for standard NWP runs. The most important namelist parameters are discussed along with recommended settings.

The main switch for activating tracer transport is $l_{transport}$ (run_nml). Except for specific idealized test cases (see Chapter 4) this switch should generally be set to .TRUE.. The namelist run_nml contains a second relevant parameter termed $n_{tracer}$ which is meant for specifying the total number of tracers that shall be advected. We note, however, that this parameter has become obsolete, except for idealized cases. In real case runs, ICON takes care of initializing the correct number of tracers based on the selected physics packages.

The namelist transport_nml contains additional parameters for selecting the transport scheme and the type of limiter. This can be done individually for each tracer, for horizontal and vertical directions.

ihadv_tracer (namelist transport_nml, list of Integer values)
Comma separated list of integer values, specifying the type of horizontal transport scheme. The $i^{th}$ entry corresponds to the $i^{th}$ tracer in ICON’s internal tracer list. Most relevant options are
3.6 Tracer Transport

1. 1st order upwind
2. MIURA (Miura (2007)-type with linear reconstruction)
3. MIURA3 (Miura (2007)-type with cubic reconstruction)
4. FFSL with quadratic or cubic reconstruction (depends on \texttt{lsq high ord (interpol nml)})
5. hybrid MIURA3/FFSL with quadratic or cubic reconstruction

Sub-cycling means that the integration from $t^n$ to $t^{n+1}$ is split into substeps to meet the stability requirements. Sub-cycling is only applied above a certain height defined by \texttt{hbot qsubjstep (nonhydrostatic nml)}. See Section 3.8.11.

FFSL and MIURA3 only differ w.r.t. the way the integration over the flux area is performed. FFSL can cope with slightly larger Courant numbers while being somewhat more expensive. Option 5 tries combine the improved stability of FFSL with the speed of MIURA3 by calling FFSL only for those edges for which the horizontal Courant number exceeds a threshold.

\texttt{ivadv_tracer (namelist transport nml, list of Integer values)}
Comma separated list of integer values, specifying the type of vertical transport scheme. The $i^{th}$ entry corresponds to the $i^{th}$ tracer in ICON’s internal tracer list. Most relevant options are

1. 1st order upwind
2. PPM

\texttt{itype_hlimit (namelist transport nml, list of Integer values)}
Comma separated list of integer values, specifying the type of horizontal limiter. The $i^{th}$ entry corresponds to the $i^{th}$ tracer in ICON’s internal tracer list. Most relevant options are

0. no limiter
1. monotonic Flux Corrected Transport (Zalesak, 1979)
2. positive definite Flux Corrected Transport

\texttt{itype_vlimit (namelist transport nml, list of Integer values)}
Comma separated list of integer values, specifying the type of vertical limiter. The $i^{th}$ entry corresponds to the $i^{th}$ tracer in ICON’s internal tracer list. Most relevant options are

0. no limiter
1. semi-monotonic reconstruction filter
2. monotonic reconstruction filter
4. positive definite Flux Corrected Transport

Example Settings for a Standard NWP Run

Valid settings for a standard NWP run with one-moment microphysics (i.e. 5 prognostic water tracers) are depicted in Figure 3.5. Currently the mapping between the $i^{th}$ entry
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! transport_nml: tracer transport --------------------------
&transport_nml
ihadv_tracer = 52, 2, 2, 2, 2 ! hor. transport selector
ivadv_tracer = 3, 3, 3, 3, 3 ! vert. transport selector
itype_hlimit = 3, 4, 4, 4, 4 ! hor. limiter
itype_vlimit = 1, 1, 1, 1, 1 ! vert. limiter
/

Figure 3.5.: Example namelist settings for tracer transport in a standard NWP run with one-moment microphysics without graupel (i.e. 5 tracers $q_v$, $q_c$, $q_i$, $q_r$, $q_s$).

in the above namelist parameters and the $i^{th}$ entry in ICON’s internal tracer list is hard coded. The order is as follows: $q_v$, $q_c$, $q_i$, $q_r$, $q_s$, i.e. the first entry relates to water vapor, the second one to cloud water, and so on. This can become unwieldy and error prone if more than a handful of tracers is used. The ART-package alleviates this problem by providing a more elaborate way of configuring tracers which is based on XML files (see Section 10.4.3). Note, however that the configuration via XML files is restricted to ART-specific tracers, only.

PPM is the method of choice in vertical direction for all tracers. In horizontal directions MIURA is used for all tracers except vapor $q_v$. A somewhat more accurate (and more expensive) scheme is selected for $q_v$ (hybrid MIURA3/FFSL). Note that sub-cycling is activated for $q_v$. This is crucial for numerical stability!

One might wonder why sub-cycling is only activated for $q_v$. The reason is that with standard NWP settings $q_v$ is the only tracer for which transport is activated all the way up to the model top where the highest wind speeds are typically encountered. For all other water tracers transport is switched off above a certain height defined by $htop$ (nonhydrostatic.nml) (typically around 18 km) such that sub-cycling is not strictly required (see Section 3.8.11).

Furthermore, note that if additional non-water tracers (e.g. purely diagnostic passive tracers or chemical tracers) are added, sub-cycling must be activated since $htop$ is only effective for water tracers.

In terms of limiters, the rule of thumb is that at least a positive definite limiter should be used for all water tracers. Otherwise numerical instabilities will occur due to negative water concentrations. For $q_v$ it is advisable to use a more stringent (albeit more expensive) monotonic limiter in order to reduce spurious condensation/evaporation emerging from nonphysical over-/undershoots in $q_v$. 

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3.7 Physics-Dynamics Coupling

3.7.1. ICON Time-Stepping

For efficiency reasons, different integration time steps are applied depending on the process under consideration. In the following, the term dynamical core refers to the numerical solution of the dry Navier-Stokes equations, while the term physics refers to the diabatic, mostly sub-grid scale, processes that have to be parameterized. In ICON, the following time steps have to be distinguished:

- $\Delta t$ the basic time step specified via namelist variable `dtime`, which is used for tracer transport, numerical diffusion and the fast-physics parameterizations.
- $\Delta \tau$ the short time step used within the dynamical core; the ratio between $\Delta t$ and $\Delta \tau$ is specified via the namelist variable `ndyn_substeps` (namelist `nonhydrostatic_nml`, number of dynamics substeps), which has a default value of 5.
- $\Delta t_{i,\text{slow physics}}$ the process dependent slow physics time steps; they should be integer multiples of $\Delta t$ and are rounded up automatically if they are not.

An illustration of the relationship between the time steps can be found in Figure 3.6. More details on the physics-dynamics coupling will be presented in Section 3.7.

ICON solves the fully compressible nonhydrostatic Navier-Stokes equations using a time stepping scheme that is explicit except for the terms describing vertical sound wave propagation. Thus, the maximum allowable time step $\Delta \tau$ for solving the momentum, continuity
3. Model Description

and thermodynamic equations is determined by the fastest wave in the system – the sound waves. As a rule of thumb, the maximum dynamics time step can be computed as

$$\Delta \tau = 1.8 \cdot 10^{-3} \frac{\Delta x}{m}$$  \hspace{1cm} (3.39)

where $\Delta x$ is the effective horizontal mesh size in meters (see Equation (2.1)). This implies that the namelist variable $dtime$ should have a value of

$$\Delta t = 9 \cdot 10^{-3} \frac{\Delta x}{m},$$

unless $ndyn\_substeps$ is set to a non-default value.

Additional time step restrictions for $\Delta t$ arise from the numerical stability of the horizontal transport scheme and the physics parameterizations, in particular due to the explicit coupling between the turbulent vertical diffusion and the surface scheme. Experience shows that $\Delta t$ should not significantly exceed 1000 s, which becomes relevant when $\Delta x$ is larger than about 125 km.

Even longer time steps than $\Delta t$ can be used for the so-called slow-physics parameterizations, i.e. radiation, convection, non-orographic gravity wave drag, and orographic gravity wave drag. These parameterizations provide tendencies to the dynamical core, allowing them to be called individually at user-specified time steps. The related namelist switches are $dt\_rad$, $dt\_conv$, $dt\_gwd$ and $dt\_sso$ in $nwp\_phy\_nml$. If the slow-physics time step is not a multiple of the advective time step, it is automatically rounded up to the next advective time step. A further recommendation is that $dt\_rad$ should be an integer multiple of $dt\_conv$, such that radiation and convection are called at the same time\(^1\). The time-splitting is schematically depicted in Figure 3.6.

For efficiency reasons, a distinction is made between so-called fast-physics processes (those whose time scale is comparable or shorter than the model time step), and slow-physics processes whose time scale is considered slow compared to the model time step. The relationship between the different time steps has already been explained in Section 3.7.1.

Fast-physics processes are calculated at every physics time step and are treated with time splitting (also known as sequential-update split) which means that (with exceptions

\(^1\)This behavior is automatically enforced in the current model version.
3.7 Physics-Dynamics Coupling

noted below) they act on an atmospheric state that has already been updated by the dynamical core, horizontal diffusion and the tracer transport scheme. Each process then sequentially updates the atmospheric variables and passes a new state to the subsequent parameterization.

The calling sequence is saturation adjustment → surface transfer scheme → land-surface scheme → boundary-layer / turbulent vertical diffusion scheme → microphysics scheme, and again saturation adjustment in order to enter the slow-physics parameterizations with an adjusted state. The exceptions from the above-mentioned sequential splitting are the surface transfer scheme and the land-surface scheme, which take the input at the ‘old’ time level because the surface variables are not updated in the dynamical core and the surface transfer coefficients and fluxes would be calculated from inconsistent time levels otherwise. The coupling strategy is schematically depicted in Figure 3.7.

Slow-physics processes are treated in a parallel-split manner, which means that they are stepped forward in time independently of each other, starting from the model state provided by the latest fast-physics process. In ICON convection, subgrid-scale cloud cover, radiation, non-orographic and orographic gravity wave drag are considered as slow processes. Typically, these processes are integrated with time steps longer than the (fast) physics time step. The slow-physics time steps can be specified by the user. The resulting slow-physics tendencies $\partial v_n/\partial t$, $\partial T/\partial t$ and $\partial q_x/\partial t$ with $x \in \{v, c, i\}$ are passed to the dynamical core and remain constant between two successive calls of the parameterization (Figure 3.7). Since ICON solves a prognostic equation for $\pi$ rather than $T$, the temperature tendencies are converted into tendencies of the Exner function, beforehand. Rather than treating the moisture tendencies as a forcing term during tracer advection, they are treated in a time-split manner and added to the updated moisture variables thereafter.

3.7.2. Isobaric vs. Isochoric Coupling Strategies

The physics-dynamics coupling in ICON differs from many existing atmospheric models in that it is performed at constant density (volume) rather than constant pressure. This is related to the fact that the total air density $\rho$ is one of the prognostic variables, whereas pressure is only diagnosed for parameterizations needing pressure as input variable. Thus, it is natural to keep $\rho$ constant in the physics-dynamics interface. As a consequence, heating rates arising from latent heat release or radiative flux divergences have to be converted into temperature changes using $c_v$, the specific heat capacity at constant volume of moist air. Some physics parameterizations inherited from hydrostatic models, in which the physics-dynamics coupling always assumes constant pressure, therefore had to be adapted appropriately.

Moreover, it is important to note that the diagnosed pressure entering into a variety of parameterizations is a hydrostatically integrated pressure rather than a nonhydrostatic pressure derived directly from the prognostic model variables$^2$. This is motivated by the fact that the pressure is generally used in physics schemes to calculate the air mass represented by a model layer, and necessitated by the fact that sound waves generated by the

$^2$Note that the (surface) pressure available for output is as well the hydrostatically integrated pressure rather than a nonhydrostatic pressure derived directly from the prognostic model variables.
Figure 3.7.: Coupling of the dynamical core and the NWP physics package. Processes declared as fast (slow) are treated in a time-split (process-split) manner.

saturation adjustment can lead to a local pressure increase with height in very extreme cases, particularly between the lowest and the second lowest model level.

Another important aspect is related to the fact that physics parameterizations traditionally work on mass points (except for three-dimensional turbulence schemes). While the conversion between different sets of thermodynamic variables is reversible except for numerical truncation errors, the interpolation between velocity points and mass points potentially induces errors. To minimize them, the velocity increments, rather than the full velocities, coming from the turbulence scheme are interpolated back to the velocity points and then added to the prognostic variable $v_n$. 

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3.8. ICON NWP-Physics in a Nutshell

An in-depth description of the physical parameterization package for NWP is beyond the scope of this document. However, the following section provides a short introduction to the available parameterizations and references for further reading.

Table 3.3 contains a summary of physical parameterizations available in ICON (NWP-mode). In what follows, an outline (executive summary) of the parameterization schemes is given.

3.8.1. Radiation

The calculation of radiation is performed with RRTM (Rapid Radiative Transfer Model, Mlawer et al., 1997). RRTM operates on 30 spectral bands with 16 bands in the long-wave spectrum and 14 bands in the shortwave spectrum. It utilizes the correlated-k method that strongly reduces computational costs with an accuracy comparable to more detailed line-by-line models.

3.8.2. Cloud Microphysics

In the exercises for Chapter 8 (see p. 240) we will investigate ICON’s physical parameterizations by means of a custom diagnostic quantity. We restrict ourselves to the cloud microphysics parameterization, where some additional background information will be of interest.

Microphysical schemes provide a closed set of equations to calculate the formation and evolution of condensed water in the atmosphere. The most simple schemes predict only the specific mass content of certain hydrometeor categories like cloud water, rain water, cloud ice and snow. This is often adequate, because it is sufficient to describe the hydrological cycle and the surface rain rate, which is the vertical flux of the mass content. Microphysical schemes of this category are called single-moment schemes.

In ICON two single-moment schemes are available, one that predicts the categories cloud water, rain water, cloud ice and snow (\texttt{inwp.gscp=1} in the namelist \texttt{nwp.phy.nml}), and the other that predicts in addition also a graupel category (\texttt{inwp.gscp=2}). Graupel forms through the collision of ice or snow particles with supercooled liquid drops, a process called riming.

Most microphysical processes depend strongly on particle size and although the mean size is usually correlated with mass content this is not always the case. Schemes that predict also the number concentrations have the advantage that they provide a size information, which is independent of the mass content. Such schemes are called double-moment schemes, because both, mass content and number concentration, are statistical moments of the particles size distribution.
### 3. Model Description

#### Table 3.3: Summary of ICON’s physics parameterizations for NWP, together with the related namelist settings (namelist nwp.phy.nml). Parameterizations which are used operationally (at 13 km horizontal grid spacing) are indicated by 

<table>
<thead>
<tr>
<th>Process</th>
<th>Scheme</th>
<th>Settings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Radiation</strong></td>
<td>RRTM (Rapid Radiative Transfer Model)</td>
<td>inwp.radiation=1</td>
</tr>
<tr>
<td></td>
<td>Mlawer et al. (1997), Barker et al. (2003)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>PSRAD</td>
<td>inwp.radiation=3</td>
</tr>
<tr>
<td></td>
<td>Pincus and Stevens (2013)</td>
<td></td>
</tr>
<tr>
<td><strong>Non-orographic gravity wave drag</strong></td>
<td>Wave dissipation at critical level</td>
<td>inwp.gwd=1</td>
</tr>
<tr>
<td></td>
<td>Orr et al. (2010)</td>
<td></td>
</tr>
<tr>
<td><strong>Sub-grid scale orographic drag</strong></td>
<td>Lott and Miller scheme</td>
<td>inwp.aso=1</td>
</tr>
<tr>
<td></td>
<td>Lott and Miller (1997)</td>
<td></td>
</tr>
<tr>
<td><strong>Cloud cover</strong></td>
<td>Diagnostic PDF</td>
<td>inwp.cldcover=1</td>
</tr>
<tr>
<td></td>
<td>M. Köhler et al. (DWD)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>All-or-nothing scheme (grid-scale clouds)</td>
<td>inwp.cldcover=5</td>
</tr>
<tr>
<td><strong>Microphysics</strong></td>
<td>Single-moment scheme</td>
<td>inwp.gscp=1, 2</td>
</tr>
<tr>
<td></td>
<td>Doms et al. (2011), Seifert (2008)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Double-moment scheme</td>
<td>inwp.gscp=4</td>
</tr>
<tr>
<td></td>
<td>Seifert and Beheng (2006)</td>
<td></td>
</tr>
<tr>
<td><strong>Convection</strong></td>
<td>Mass-flux shallow and deep</td>
<td>inwp.convection=1</td>
</tr>
<tr>
<td></td>
<td>Tiedtke (1989), Bechtold et al. (2008)</td>
<td></td>
</tr>
<tr>
<td><strong>Turbulent transfer</strong></td>
<td>Prognostic TKE (COSMO)</td>
<td>inwp.turb=1</td>
</tr>
<tr>
<td></td>
<td>Raschendorfer (2001)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>EDMF-DualM (Eddy-Diffusivity/Mass-Flux)</td>
<td>inwp.turb=3</td>
</tr>
<tr>
<td></td>
<td>Köhler et al. (2011), Neggers et al. (2009)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3D Smagorinsky diffusion (for LES)</td>
<td>inwp.turb=5</td>
</tr>
<tr>
<td><strong>Land</strong></td>
<td>Tiled TERRA</td>
<td>inwp.surface=1</td>
</tr>
<tr>
<td></td>
<td>Schrodin and Heise (2001), Schulz et al. (2016)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Flake: Mironov (2008)</td>
<td>llake=.TRUE.</td>
</tr>
<tr>
<td></td>
<td>Sea-ice: Mironov et al. (2012)</td>
<td>lseaice=.TRUE.</td>
</tr>
</tbody>
</table>
ICON does also provide a double-moment microphysics scheme (\texttt{inwp.gscp=4}), which predicts the specific mass and number concentrations of cloud water, rain water, cloud ice, snow, graupel and hail. This scheme is most suitable at convection-permitting or convection-resolving scales, i.e., mesh sizes of 3 km and finer. Only on such fine meshes the dynamics is able to resolve the convective updrafts in which graupel and hail form. On coarser grids the use of the double-moment scheme is not recommended.

To predict the evolution of the number concentrations the double-moment scheme includes various parameterizations of nucleation processes and all relevant microphysical interactions between these hydrometeor categories. Currently all choices regarding, e.g., cloud condensation and ice nuclei, particle geometries and fall speeds etc. have to be set in the code itself and can not be chosen via the ICON namelist.

### 3.8.3. Cumulus Convection

D. Klocke, DWD Physical Processes Division

Convection is an important process in the atmosphere by contributing to forming the large-scale circulation to local heavy precipitation through thunderstorms. Parameterizations of atmospheric moist convection provide the effect of an ensemble of sub-grid convective clouds on the model column as a function of grid-scale variables. The schemes vertically mix heat, moisture and momentum. They convert available potential energy into kinetic energy and produce precipitation as a result of atmospheric instability.

Three steps are taken. First, it is determined if the grid-scale conditions allow for the occurrence of convection in the column, and a decision is taken if convection is triggered. In the second step, the tendencies of heat, moisture and momentum changes are determined with a cloud model, which represents an ascending parcel and its interactions with the environment. Finally the closure decides on the strength of the convection by determining the amount of energy to be converted, which is linked to precipitation amount generated by the convection scheme.

In ICON a bulk mass flux convection scheme is available \texttt{inwp.convection=1} (in the namelist \texttt{nwp.phy.nml}), which treats three convective cloud types. Only one type - shallow, mid-level or deep convection - can exist at a time in a column, which is decided upon by the trigger function. All three types of convection use a cloud model representing an ascending plume mixing with its environmental air. The cloud base mass flux closure differs between the three convection types, with a CAPE (convective available potential energy) based closure for deep convection, a boundary layer equilibrium closure for shallow convection, and a large-scale omega (vertical velocity in pressure coordinates) based closure for mid-level convection.

The full convection scheme can generally be used for horizontal grids coarser than 5 km, as some resolution dependent adjustments are implemented for grid spacings smaller than 20 km. For convection permitting simulations (1 – 3 km horizontal grid spacing) the largest convective clouds can be resolved by the model and the parameterization parts treating deep and mid-level convection can be switched off (\texttt{lshallowconv_only=.TRUE.} in the namelist \texttt{nwp.phy.nml}).
The implemented scheme represents a spin-off of the Tiedtke-Bechtold convection scheme used in the IFS model. For further reading we refer to Bechtold et al. (2008), Tiedtke (1989), Bechtold (2017), ECMWF (2017).

### 3.8.4. Cloud Cover

To prepare optical properties of clouds for the radiative transfer it is necessary to determine the best estimate of cloud cover, cloud water and cloud ice as well as the precipitation quantities, such as snow, rain and graupel if those are required by the radiation calculation. Note that there are various assumptions in the ICON model on the subgrid distribution of water, such as the up/down/subsidence regions in convection, a uniform distribution in microphysics, and a Gaussian distribution in turbulence.

The aim of the diagnostic cloud cover scheme is to combine information from the different parameterizations mentioned above: turbulence, convection and microphysics. The turbulence scheme provides the sub-grid variability of water due to turbulent motions, the convection scheme detrains cloud into the anvil and the microphysics scheme describes the supersaturation due to ice, in other words, the distribution between ice and vapor in cold situations.

The turbulent variability of water is at the moment prescribed by using a top-hat total water (the sum of water vapor $q_v$, cloud water $q_c$, and cloud ice $q_i$) distribution with a fixed width of 10% of total water. Work is in progress to replace this crude assumption with the total water variance from the turbulence scheme.

The split between water vapor and cloud ice as determined from the microphysics scheme is replicated in the diagnostic cloud scheme for the turbulent component of ice clouds.

The convective anvil is calculated by writing an equation for the evolution of cloud cover that depends on the detrainment of volume (from the convection scheme) and a decay term with a fixed decay time-scale (taken as 30 min). The diagnostic assumption means that we can neglect the tendency term on cloud cover so that we arrive simply at the diagnostic anvil cloud cover that is purely a function of detrainment and decay time-scale. The liquid and ice cloud water is taken also from the convective updraft properties.

In the end the turbulent and convective clouds are combined with a simple maximum function.

To emphasize, the cloud cover scheme takes into account the subgrid variability of water and therefore the associated distribution in water vapor, cloud liquid water and cloud ice (optional 3D diagnostic output variables $\texttt{tot\_qv\_dia}$, $\texttt{tot\_qc\_dia}$, $\texttt{tot\_qi\_dia}$ and their vertically integrated counterparts $\texttt{tqv\_dia}$, $\texttt{tqc\_dia}$, $\texttt{tqi\_dia}$). They are not equal to their prognostic grid-scale equivalents (standard output variables $q_v, q_c, q_i$), yet the sum of all three water quantities is kept the same. This cloud information is then passed to the radiation, where additional assumptions are made on the vertical overlap of clouds.
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### 3.8.5. Turbulent Diffusion

Section authors
M. Köhler and M. Raschendorfer,
DWD Physical Processes Division

**TKE Scheme for Turbulence.**

The TKE turbulence scheme consists of two components: one describing the free troposphere, and the other for the surface layer.

**TURBDIFF.** The turbulence scheme TURBDIFF developed by Raschendorfer (2001) is based on a 2nd-order closure on level 2.5 according to Mellor and Yamada (1982) (MY-scheme). In this scheme, all pressure correlation terms and dissipation terms, being present in the system of 2nd-order equations for all the turbulent (co-)variances that can be built from the dynamically active prognostic model variables, are expressed by the standard closure assumptions according to Rotta (1951a,b) and Kolmogorov (1968) valid for quasi-isotropic turbulence. These dynamically active model variables are the horizontal wind components u and v, vertical wind speed w, a variable related to inner energy (like absolute temperature T, potential temperature θ or virtual potential temperature θ_v), specific humidity q_v and at least one cloud water variable q_c (which is a mass fraction and may be split into liquid q_l and frozen q_i cloud water).

Among these 2nd-order moments, only the trace of the turbulent stress tensor, which is twice the Turbulent Kinetic Energy (TKE), is described by a prognostic equation. Each of the remaining 2nd-order equations (for the elements of the remaining trace-less stress tensor and for the other 2nd-order moments) is simplified as diagnostic source term equilibrium being a linear equation in terms of the governing statistical moments.

Further, correlations between any model variable and source terms of scalar model variables are neglected in these equations. However, by choosing quasi-conserved scalar variables (total water content q_t = q_v + q_c and liquid-water potential temperature θ_l = θ - \(\frac{L_v}{c_p} q_c\)), these correlations are taken into account implicitly as far as local condensation and evaporation within liquid non-precipitating clouds is concerned. Hence, TURBDIFF is a moist turbulence scheme, which includes the effect of these sub-grid scale phase transitions. The required conversion of turbulent fluxes of these conserved variables into those of absolute temperature, specific humidity and liquid water content is performed by means of turbulent saturation adjustment, assuming a Gaussian distribution-function for local saturation-deficiency according to Sommeria and Deardorff (1977).

Finally, application of the horizontal boundary layer approximation reduces the linear system of diagnostic 2nd-order equilibrium equations to a single column scheme with only two equations for two diffusion coefficients (one for horizontal wind components and another for scalar variables), which both are proportional to the square root of TKE and an integral turbulent length scale. This length-scale rises with height above ground according to Blackadar (1962) with a further limitation related to the horizontal grid scale. The desired vertical turbulent fluxes of any prognostic variable can then be calculated by
multiplying the (negative) vertical gradient of the latter with the associated diffusion
coefficient.

One main extension of TURBDIFF (compared with a moist MY-scheme) is the formal
separation of turbulence from a possible non-turbulent part of the subgrid-scale energy
spectrum. This separation is related to additional scale-interaction terms in the prognostic
TKE equation, which describe additional shear production of TKE by the action of other
non-turbulent sub-grid-scale flow patterns (such as wakes generated by sub-grid-scale orog-
raphy, convective currents or separated horizontal circulations). Through this formalism,
the scheme describes Separated Turbulence Interacting with non-turbulent Circulations
(STIC), which allows for a consistent application of turbulence closure assumptions, even
though other sub-grid-scale processes may be dominant within a grid cell. Due to this
extension, the scheme is applicable also above the boundary layer and for very stable
stratification. The Eddy Dissipation Rate (EDR) calculated by TURBDIFF can even be
used to forecast the intensity of Clear Air Turbulence (CAT).

**TURBTRAN.** The turbulence scheme TURBDIFF is closely related to the scheme
TURBTRAN developed by Raschendorfer (2001) for the surface-to-atmosphere transfer
(SAT), which calculates transport resistances for fluxes of prognostic model variables at
the surface of the Earth. Figure 3.8 illustrates the corresponding sub layers of the surface
layer. In TURBTRAN, a constant flux approximation is applied to the sum of turbulent
and laminar vertical fluxes within the transfer layer (between the rigid surface and the
lowest atmospheric main level of the model). By application of the turbulence scheme at
the top of the lowest atmospheric layer as well as the bottom of this layer (which is the top
of the near surface roughness layer being intersected by roughness elements), a vertical
interpolation function for the turbulent diffusion coefficient is derived between these two
levels and is extrapolated down to the rigid surface. With this preparation, a vertical in-
tegration of the flux gradient representation across the transfer layer provides the desired
transport resistances for the final bulk representation of SAT fluxes. With this formula-
tion, scale interaction terms considered through STIC in the turbulence scheme also affect
the transfer resistances; and hence, some additional mixing is automatically introduced
for very large bulk Richardson numbers, as soon as non-turbulent sub-grid-scale motions
are present.

Further, with the described procedure, the determination of a specific roughness length
for scalars is substituted by a direct calculation of the partial resistance of scalar transfer
through the laminar layer and the roughness layer. This partial resistance is dependent on
the near-surface model variables, the aerodynamic roughness length and the Surface Area
Index (SAI), which is a measure of the surface area enlargement by land use.

TURBDIFF and TURBTRAN are the default schemes for atmospheric turbulence and
SAT, respectively, in ICON, and they correspond to **inwp_turb=1** (namelist **nwp.phy.nml**).
While TURBTRAN provides the transfer resistances for scalars and horizontal wind com-
ponents, the final surface fluxes of water vapor and sensible heat are determined in TERRA
as a result of updated surface values for $q_v$ and (in case of the upcoming implicit treatment
of surface temperature) also for $T$. Based on these surface fluxes, an implicit equation for
vertical diffusion is solved as a final part of TURBDIFF. In this procedure also horizontal
momentum and TKE is included, while for these 3 quantities the respective surface con-
centration is used as a lower boundary condition. Currently, a lower zero-concentration
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condition is applied for cloud-water and ice, which is always related to a downward flux for these quantities. For vertical diffusion of passive tracers, the diffusion coefficient for scalars is applied, and the lower boundary condition can be specified individually. Although the effect of local condensation and evaporation is considered in the solution of 2nd order equations, and hence, can amplify the intensity of turbulent vertical mixing, the direct effect of these additional thermodynamic source-terms in the grid-scale budgets of heat, water vapor and liquid water is not yet considered.

In the namelist `turbdiff.nml` several parameters or selectors for optional calculations related to both schemes can be specified. Through this, TURBDIFF can also be configured as a 3D-turbulence scheme, calculating additionally horizontal shear and providing also horizontal diffusion coefficients.

DualM EDMF for Turbulence and Shallow Convection.

The Eddy Diffusivity Mass Flux (EDMF) approach - operational at ECMWF (Köhler et al., 2011) - is based on the decomposition of the turbulent transports proposed by Siebesma and Cuijpers (1995) into eddy diffusivity and mass-flux components. This is based on the idea of unifying the eddy diffusivity and mass-flux concepts that are used in many NWP and climate models within one unified solver. When generalizing to multiple updrafts $M_i$ one arrives at the following equation for the flux of a scalar quantity $\phi$:

$$ \overline{w'\phi'} = -K \frac{\partial \overline{\phi}}{\partial z} + \sum_i M_i (\overline{\phi_i} - \overline{\phi}). $$

Here, the mass flux is $M_i = a_i (\overline{w_i} - \overline{w})$ and $K$ is the diffusion coefficient. The averaging operators $\overline{\phi_i}$ and $\overline{\phi}$ act on the updraft and environment fractions, respectively.
To arrive at this equation two assumptions are applied: (i) updraft fraction is small \( a \ll 1 \) and (ii) the flux within the environment \( \overline{w'} \overline{\varphi'} \) can be approximated by K-diffusion.

The DualM framework by Neggers et al. (2009) postulates that two mass-fluxes are sufficient to treat the transition from a dry boundary layer to stratocumulus and shallow cumulus. In particular one dry mass-flux stops at cloud base, while the second moist mass-flux reaches to cloud top. The continuous area partitioning between the dry and moist updraft is a function of moist convective inhibition above the mixed layer top. Updraft initialization is a function of the updraft area fraction and is therefore consistent with the updraft definition. It is argued that the model complexity thus enhanced is sufficient to allow reproduction of various phenomena involved in the cloudsubcloud coupling, namely (i) dry countergradient transport within the mixed layer that is independent of the moist updraft, (ii) soft triggering of moist convective flux throughout the boundary layer, and (iii) a smooth response to smoothly varying forcings, including the reproduction of gradual transitions to and from shallow cumulus convection.

**Smagorinsky-Lilly for LES Application**

The 3D sub-grid model of Smagorinsky (1963) with the stability correction of Lilly (1962) is implemented for LES applications. This scheme writes the eddy viscosity \( K_m \) as

\[
K_m = (C_s \Delta)^2 |S| C_B,
\]

with the Smagorinsky constant \( C_s \), the filter width \( \Delta \), the norm of the strain rate tensor \( |S| \) and the stability correction factor \( C_B \).

The filter width is taken to be \( \Delta = (\Delta_x \Delta_y \Delta_z)^{\frac{1}{3}} \). The strain rate tensor is defined as

\[
S_{ij} = \frac{1}{2} \left( \frac{\partial u_i}{\partial x_j} + \frac{\partial u_j}{\partial x_i} \right),
\]

whose calculation requires special care due to the triangular grid in ICON. A metric correction has been developed that treats the horizontal gradients over a sloped orography given a terrain-following coordinate correctly. The norm of \( S_{ij} \) is

\[
|S| = \sqrt{2 S_{ij} S_{ij}}.
\]

The stability correction factor \( C_B \) is given by

\[
C_B = (1 - \text{Ri} / \text{Pr}_t)^{\frac{1}{2}}
\]

with the gradient Richardson number \( \text{Ri} = \frac{N^2}{|S|} \) and the buoyancy frequency \( N^2 = \frac{g}{\theta_0} \frac{\partial \theta}{\partial z} \).

The Smagorinsky constant usually has the value of \( C_s = 0.1 - 0.2 \). In the ICON implementation \( C_s \) is set by default to \textit{smag.constant}=0.23 (namelist les.nml) and the Prandtl number \( \text{Pr}_t \) to \textit{turb.prandtl}=0.333 (namelist les.nml).
3.8.6. Sub-grid scale orographic drag

ICON treats the entire sub-grid scale orographic drag with one model based on the work of Lott and Miller (1997). Other NWP models treat scales smaller than $\sim 5$ km with different ansatzes, for example with the help of turbulent orographic form drag formulations of the use of an effective roughness length that determines the conductivity of the surface for momentum fluxes between the atmosphere and the Earth.

The motivation for the implementation of a sub-grid scale orographic drag model into ICON traces back to experience with the COSMO model. When the 7-km EU domain of the operational COSMO model at DWD was expanded in order to cover almost all Europe (see Schulz, 2006), it turned out that the surface pressure in the model forecasts became systematically biased. In particular, in wintertime high pressure systems of the model atmosphere tended to develop a positive pressure bias, by 1 to 2 hPa after 48 h, low pressure systems a negative bias (“highs too high, lows too low”). At the same time the wind speed tended to be overestimated by up to 1 m s$^{-1}$ throughout the entire troposphere. The wind direction near the surface showed a positive bias.

The combination of these deficiencies led to the hypothesis that in the model there is too little surface drag, causing an underestimation of the cross-isobar flow in the planetary boundary layer. Consequently, the solution would be to increase the surface drag in the model. This may be accomplished, for instance, by introducing an envelope orography (Wallace et al., 1983, Tibaldi, 1986), but this has unfavorable effects, e.g., for the simulated precipitation. Another option is the inclusion of sub-grid scale orographic (SSO) effects, which were neglected in the COSMO model before. The SSO scheme by Lott and Miller (1997) was selected for this purpose. Its implementation in the COSMO-EU model is described in Schulz (2008), and was later transferred to the ICON model.

The SSO scheme by Lott and Miller (1997) deals explicitly with a low-level flow which is blocked when the sub-grid scale orography is sufficiently high. For this blocked flow separation occurs at the mountain flanks, resulting in a form drag. The upper part of the low-level flow is led over the orography, while generating gravity waves$^3$. In order to describe the low-level flow behavior in the SSO scheme a non-dimensional height $H_n$ of the sub-grid scale mountain is introduced

$$H_n = \frac{NH}{|U|} = Fr^{-1},$$

where $H$ is the maximum height of the mountain, $|U|$ is the wind speed and $N$ is the Brunt-Väisälä frequency of the incident flow. The latter is defined by

$$N = \sqrt{\frac{g \partial \theta}{\partial z}},$$

$^3$Propagating, coherent structures consisting of buoyancy and inertial oscillations.
3. Model Description

Figure 3.9: Left: Schematic illustration of the two elements of sub-grid scale orographic drag: First, the low-level blocking, where the horizontal flow (orange) is forced to flow around the sub-grid scale orography (SSO, colored gray). Second, the gravity wave drag on the flow forced to overflow the SSO, and on the flow at higher altitudes, where the radiated gravity waves (white-blue) break. (Following Figure 1 in Lott and Miller (1997).) Right: Schematic illustration of the non-orographic gravity wave drag.

where $\theta$ is the potential temperature, $g$ the acceleration of gravity and $z$ the height coordinate. $H_n$ may be also regarded as an inverse Froude number $Fr^{-1}$ of the system “flow round a mountain”.

A small $H_n$ means that there is an unblocked regime, all the flow goes over the mountain and gravity waves (GWs) are forced by the vertical motion of the fluid. A large $H_n$ means that there is a blocked regime, the vertical motion of the fluid is limited and part of the low-level flow goes around the mountain. The SSO scheme requires four external parameters, which are the standard deviation \texttt{SSO\_STDH}, the anisotropy \texttt{SSO\_GAMMA}, the slope \texttt{SSO\_SIGMA} and the geographical orientation \texttt{SSO\_THETA} of the sub-grid scale orography. Following Baines and Palmer (1990) these are computed by ExtPar (see Section 2.4.1) from the GLOBE data set (GLOBE-Task-Team, 1999), which has a resolution of approximately 1 km.

Four tuning parameters in the namelist \texttt{nwp\_tuning.nml} control the SSO scheme (see Figure 3.9). First, the critical Froude number $Fr_c = \texttt{tune\_gfrcrit}$, which has a twofold effect. The larger its value the higher the likelihood for low-level blocking to occur, since it is activated only where $Fr < Fr_c$. Conversely, where $Fr > Fr_c$ all flow goes over the mountain and the entire stress is associated with GW radiation. In addition, if blocking is active, $Fr_c$ controls the thickness of the blocking layer relative to the mountain height (the larger $Fr_c$ the larger the layer thickness). Its default value is 0.4. Operational simulations with horizontal mesh sizes of 13 km and 40 km use the values 0.333 and 0.425, respectively. Second and third, the magnitudes of the SSO drag and the GW drag are directly proportional to the parameters $\texttt{tune\_gkwake}$ and $\texttt{tune\_gkdrag}$, respectively. The default values are 1.5 and 0.075. Different from this, operational simulations with 13 km horizontal mesh size use $\texttt{tune\_gkwake} = 1.8$ and $\texttt{tune\_gkdrag} = 0.09$. Finally, the critical Richardson number $Ri_c = \texttt{tune\_grcrit}$ is a control parameter for the onset of GW
breaking. If the Richardson number of the resolved atmospheric state plus the unresolved GWs

\[ Ri = \frac{N_{tot}^2}{|\partial u_{tot}/\partial z|^2}, \]

where \( u \) denotes the horizontal wind vector, falls below \( Ri_c \), the flow configuration becomes unstable and the GWs break (partly). This process is accompanied by a drag effect on the resolved horizontal flow. Combined with the rule of thumb that \( Ri \) increases with height if GWs are present, this means that the larger the value of \( Ri_c \) the lower the altitude where the radiated GWs tend to break and exert a drag. The default value is 0.25.

In order to assist numerical stability, the drag computed by the scheme is limited to a height-dependent threshold (see Section 3.8.7 for some further details).

### 3.8.7. Non-orographic gravity wave drag

All kinds of (synoptic-scale) atmospheric flow structures (e.g., fronts, convection, jet streams) can develop imbalances that force air parcels to oscillate (vertically) and radiate gravity waves (GWs). The interaction of these non-orographically forced GWs with the atmospheric background flow is assumed to be significant in the middle and upper atmosphere, and is consequently of interest for models that cover this region (e.g., the operational global ICON configuration with its model top at 75 km). If the synoptic scale flow (be it resolved or unresolved by the model) would force GWs, whose horizontal and vertical wave lengths would be smaller than the horizontal and vertical grid mesh sizes, they cannot be resolved by the model. But yet this unresolved part of the GW spectrum could have a significant impact on the resolved flow, and is therefore parameterized (see Figure 3.9).

The parameterization implemented in ICON follows the ansatz of Scinocca (2003) and McLandress and Scinocca (2005), which in turn is based on the work of Warner and McIntyre (1996) to which the simplifying assumption of a hydrostatic, non-rotational atmosphere has been applied. The parameters of this scheme have been optimized following Ern et al. (2006). Details can be found in Orr et al. (2010).

The mechanisms of non-orographic GW forcing can be relatively complex and are not completely understood. For this reason the parameterization considers a statistical mean of all sources of (upward propagating) GWs and represents them by prescribing GW-borne vertical fluxes of horizontal momentum at a launch level somewhere in the upper troposphere. The magnitude of the integral of the momentum fluxes over the GW spectrum is assumed to be constant (i.e. independent of the geographic location and time), and can be expressed by

\[ \rho F (\cos \varphi_i, \sin \varphi_i)\big|_{z=z_0} = \rho_0 F_0 (\cos \varphi_i, \sin \varphi_i) = \rho \bar{w}'\bar{u}'\big|_{z=z_0} = \rho \bar{w}'\bar{u}' (\cos \varphi_i, \sin \varphi_i)\big|_{z=z_0}, \]
where \( \rho \) is the air density, \( z_0 \) denotes the launch level, \( u' \) and \( u' \) are GW-borne fluctuations of the vertical and horizontal wind components and \( \langle \cdot \rangle \) stands for a spatial average over the grid cell size. The azimuthal angles relative to the West-East direction \( \varphi_i \in \{0^\circ\) (East), \( 90^\circ\) (North), \( 180^\circ\) (West), \( 270^\circ\) (South)\} are four horizontal directions of horizontal momentum sampled by the scheme. The magnitude \( \rho_0 F_0 = \text{tune}\_gflxlaun \) can be set in namelist \text{nwp\_tuning\_nml}. Its default value is 0.0025 Pa.

Given the atmospheric state in a grid cell column and the spectral distribution of \( \rho F \) at the launch level, standard GW theories can be used, to integrate \( \rho F \) vertically. The drag exerted on the resolved horizontal flow is then defined by the vertical divergence of the momentum flux

\[
\left( \frac{\partial u}{\partial t} \right)_{\text{GW drag}} = \left[ \frac{\partial (u,v)}{\partial t} \right]_{\text{GW drag}} = -\frac{1}{\rho} \rho F_{\text{zonal}} \frac{\partial}{\partial z},
\]

(3.40)

Here, \( \langle \cdot \rangle \) denotes the integral over the GW spectrum. The right-hand side becomes non-zero typically at altitudes, where the GWs break. Note, if the magnitude of the drag computed by this scheme exceeds a prescribed, height-dependent threshold value \( \| \langle \partial u/\partial t \rangle_{\text{GW drag}} \|_{\text{max}}(z) \), it is reduced to this value. This measure is to counteract numerical instabilities caused by a too strong increase of the wind magnitude. It also applies to wind tendencies from the SSO scheme described in Section 3.8.6.

The processes treated by this parameterization and the one described in Section 3.8.6 are assumed to be irreversible and as such provide a source of heat. The corresponding temperature tendency is derived from Gibbs equation (e.g. Falk, 1990)

\[
\delta e = u \cdot \delta p + T \delta s + \cdots,
\]

(3.41)

which is one possible formulation of a fundamental relation \( F(e, p, s, \ldots) = 0 \) defined to hold between energy density \( e \), momentum density \( p \), heat density \( s \) and the densities of all other extensive state variables contained by the system\(^4\). In addition, \( \delta \) denotes the difference between two consecutive states, typically expanded by \( \delta(\cdot) = \partial(\cdot)/\partial t \delta t \), where \( \delta t \) is a time step. For the extensive variables on the right-hand side of Eq. (3.41) explicit budget equations exist in the model in one form or another. In contrast, energy is not explicitly budgeted, but its change is determined implicitly through Gibbs equation. Now, energy shall be a conserved quantity. Given that we know \( \delta p \) from the drag (3.40), whereas the heat change \( \delta s \) is still unspecified, one possibility to ensure energy conservation is to equate the right-hand side of Eq. (3.41) to zero by\(^5\)

\[
\delta s = -\frac{u \cdot \delta p + \cdots}{T}.
\]

(3.42)

\(^4\)Extensive variables depend on the amount of substance, intensive variables do not. Example: if you merge two identical systems to one single larger system, its velocity is still the same as with the two original systems, whereas its momentum is twice as large, its temperature is still the same, whereas its amount of heat is twice as large etc.

\(^5\)Strictly speaking, this holds only, if we would give up the (arbitrary) distinction in what we call “model dynamics” and “model physics” and treat both in a uniform, consistent way. In addition, time integration would practically have to be limited to an explicit Euler method, since higher-order schemes mix up different states, which challenges energy conservation and the satisfaction of the integrability conditions on Gibbs equation, known as the Maxwell relations between the equations of state. Beyond that implicit methods violate causality. Of course, all that are unrealizable requirements on an operational model for a long time to come.
3. Model Description

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Combined with the equations of state (and some simplifications) this can be formulated as a temperature tendency, which reads

\[
\left( \frac{\partial T}{\partial t} \right)_{\text{GW drag}} = -\frac{u}{c_v} \cdot \left( \frac{\partial u}{\partial t} \right)_{\text{GW drag}} = -\frac{1}{c_v} \left[ u \left( \frac{\partial u}{\partial t} \right)_{\text{GW drag}} + v \left( \frac{\partial v}{\partial t} \right)_{\text{GW drag}} \right]
\]

for the drag (3.40). However, one should keep in mind that the ansatz (3.42) is a “brute force method” applied for convenience. If we consider only state changes due to the parameterization, the consequence of Eq. (3.42) is that the amount of energy contained in a cell is “frozen”. This is because Eq. (3.42) does not only suppress energy sources, but also energy fluxes between neighboring cells that could result from the parameterized process. Fluxes, however, are generally not in conflict with conservation principles, so Eq. (3.42) might be regarded as an unrealistic simplification. A more gentle approach would follow from the general treatment of Gibbs equation in irreversible thermodynamics (e.g. Herbert, 1975, Sievers, 1982). Given that the change of momentum follows from a flux divergence \( \delta p = -\partial F/\partial z \delta t \) (compare Eq. (3.40)), Eq. (3.41) can be formulated as

\[
\delta e = -\delta t u \cdot \frac{\partial F}{\partial z} + \cdots = -\delta t \left[ \frac{\partial (u \cdot F)}{\partial z} - F \cdot \frac{\partial u}{\partial z} \right] + \cdots,
\]

to separate the energy flux \( u \cdot F \) from the “actual” source term, which would then be added to the heat sources, to satisfy energy conservation

\[
\delta s = -\delta t \frac{F}{T} \cdot \frac{\partial u}{\partial z} + \cdots.
\]  

(3.43)

However, with respect to computational costs, the variant (3.43) is typically significantly more expensive than Eq. (3.42)), which is the reason, why the latter is preferred.

The preceding considerations were guided by the focus on the energy and the demand for its conservation (the 1\textsuperscript{st} law of thermodynamics). Another aspect comes into play, if we consider heat and our experimental experience that it is not conserved, but can be produced (the 2\textsuperscript{nd} law of thermodynamics). If the drag (3.40) would be the only contribution to the right-hand side of Eq. (3.41) in addition to the contribution of heat, the 2\textsuperscript{nd} law would require \( \delta s = -\delta t/T \cdot F \cdot \partial u/\partial z \geq 0 \). This could only be satisfied, if \( F \propto -\partial u/\partial z \) (the typical gradient ansatz, we know from the formulation of friction, for instance). But that changes, if we complement Gibbs equation by the contributions from the gravity waves, turbulence etc., and allow for cross effects. Nevertheless, we should not expect the physics parameterizations to satisfy the 2\textsuperscript{nd} law, neither locally nor globally. On the one hand the 2\textsuperscript{nd} law is usually not one of the guiding principles in their development, on the other hand the different treatment of what we call “model dynamics” and “model physics”, albeit necessary for computational efficiency, challenges both energy conservation and the indestructibleness of heat.
Lakes significantly affect the structure and the transport properties of the atmospheric boundary layer. The interaction of the atmosphere with the underlying surface strongly depends on the surface temperature.

In numerical weather prediction (NWP), a simplified approach is often taken that amounts to keeping the water surface temperature constant over the entire forecast period. This approach is to some extent justified for ocean and seas. It is hardly applicable to lakes where diurnal variations of the water surface temperature reach several degrees. The situation is even more grave for frozen lakes as the diurnal variations of the ice surface temperature may exceed ten degrees.

Initialization of the NWP model grid boxes that contain water bodies also presents considerable difficulties. When no observational data for some grid boxes are available, those grid boxes are initialized by means of interpolation between the nearest grid-boxes for which the water surface temperature is known (the interpolation procedure may account for some other variables, e.g., two-meter temperature over land). Such procedure is acceptable for sea points where large horizontal gradients of the water surface temperature are comparatively rare, but it is hardly suitable for lakes. Lakes are enclosed water bodies of a comparatively small horizontal extent. The lake surface temperature has little to do with the surface temperature obtained by means of interpolation between the alien water bodies.

In NWP, the lake surface temperature (i.e., the surface temperature of lake water or lake ice) is a major concern. It is this variable that communicates information between the lake and the atmosphere, whereas details of the vertical temperature distribution (e.g., the temperature near the lake bottom) are of minor importance. Therefore, simplified lake models (parameterization schemes), whose major task is to predict the lake surface temperature, the lake freezing and the ice break-up, should be sufficient for NWP and related applications.

The NWP model ICON (as well as COSMO) utilizes the lake parameterization scheme FLake \cite{Mironov2008, Mironov2010, Mironov2012}. FLake is based on a two-layer parametric representation of the evolving temperature profile. The structure of the stratified layer between the upper mixed layer and the basin bottom, the lake thermocline, is described using the concept of self-similarity (assumed shape) of the temperature-depth curve. The same concept is used to describe the temperature structure of the thermally active upper layer of bottom sediments and of the ice and snow cover. In this way, the problem of solving partial differential equations (in depth and time) for the temperature and turbulence quantities is reduced to solving ordinary differential equations (in time only) for the time-dependent parameters that specify the temperature profile.

The approach is based on what is actually “verifiable empiricism”. However, it still incorporates much of the essential physics and offers a very good compromise between physical realism and computational economy. FLake incorporates the heat budget equations for the four layers in question, viz., snow, ice, water and bottom sediments, developed with due
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regard for the volumetric character of the solar radiation heating. An entrainment equation is used to compute the depth of a convectively-mixed layer, and a relaxation-type equation is used to compute the wind-mixed layer depth in stable and neutral stratification. Simple thermodynamic arguments are invoked to develop the evolution equations for the ice and snow depths.

Empirical constants and parameters of FLake are estimated, using independent empirical and numerical data. They should not be re-evaluated when the scheme is applied to a particular lake. In this way, the scheme does not require re-tuning, a procedure that may improve an agreement with a limited amount of data but should generally be avoided. Further information about FLake can be found at http://lakemodel.net.

FLake is activated within ICON if the namelist parameter l1ake (lnd_nml) is set .TRUE., which is the default operational setting at DWD.

FLake requires two external parameter fields. These are the fields of lake fraction (area fraction of a given numerical-model grid box covered by the lake water) and of lake depth. These external parameter fields are generated with the ExtPar software (see Section 2.4) using the Global Lake Database (Kourzeneva, 2010, Kourzeneva et al., 2012, Choulga et al., 2014).

ICON makes use of a tile approach to compute the grid-box mean values of temperature and humidity (and of other scalars) and the grid-box mean fluxes of momentum and scalars. FLake is applied to the ICON grid boxes whose lake fraction exceeds a threshold value; otherwise the effect of sub-grid scale lakes is ignored. Currently, the value of 0.05 is used (see the namelist variable frlake_thrthld (lnd_nml)).

In the current ICON configuration, the lake depth is limited to 50 m. For deep lakes, the abyssal layer is ignored, a “false bottom” is set at a depth of 50 m, and the bottom heat flux is set to zero. The bottom sediment module is switched off, and the heat flux at the water-bottom sediment interface (or at false bottom) is set to zero. The setting lflk_botsed_use=.FALSE. is hard-coded in mo_data_flake.f90.

Snow above the lake ice is not considered explicitly. The effect of snow is accounted for in an implicit manner through the temperature dependence of the ice surface albedo with respect to solar radiation Mironov et al. (2012). There is no logical switch to deactivate the snow module of FLake. It is sufficient to set the rate of snow accumulation to zero (hard-coded in mo_flake.f90). Without explicit snow layer of the lake ice, the snow depth over lakes is set to zero and the snow surface temperature is set equal to the ice surface temperature.

The attenuation coefficient of lake water with respect to solar radiation is currently set to a default “reference” value for all lakes handled by ICON. It would be advantageous to specify the attenuation coefficient as a global external parameter field. This can be done in the future as the information about the optical properties of lakes becomes available (not the case at the time being).

Generally, no observational data are assimilated into FLake, i.e., the evolution of the lake temperature, the lake freeze-up, and break-up of ice occur freely during the ICON runs. An exception are the Laurentian Great Lakes of North America. Over the Laurentian Great Lakes, the observation data on the ice fraction (provided by the ICON surface
The ice thickness, the ice surface temperature, and (as needed) the water temperature. See the subroutine `flake_init` in `mo_flake.f90` for details. The use of the ice-fraction data over Great Lakes is controlled by the namelist parameter `use_lakeiceana (initicon_nml)`. Finally, a word of caution is in order. Running ICON with the lake parameterization scheme switched off (`llake=.FALSE.`) is not recommended as this configuration has never been comprehensively tested at DWD.

### 3.8.9. Sea-Ice Parameterization Scheme

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A major task of the sea-ice parameterization scheme for NWP is to predict the existence of ice within a given atmospheric-model grid box and the ice surface temperature. The sea-ice scheme used within ICON NWP accounts for thermodynamic processes only, i.e., no ice rheology is considered (cf. the sea-ice scheme for climate modeling). The horizontal distribution of the ice cover, i.e., the fractional area coverage of sea ice within a given grid box, is governed by the data assimilation scheme. A detailed description of the sea-ice scheme for ICON NWP is given in Mironov et al. (2012), where a systematic derivation of governing equations, an extensive discussion of various parameterization assumptions and of the scheme disposable parameters, and references to relevant publications can be found. Further comments can be found directly in the code, see the module `src/ind_phy_schemes/mo_seaice_nwp.f90`.

A distinguishing feature of the ICON NWP sea-ice scheme is the treatment of the heat transfer through the ice. As different from many other sea-ice schemes that solve the heat transfer equation on a finite difference grid, the present scheme uses the integral, or bulk, approach (cf. the lake parameterization scheme FLake, Section 3.8.8). It is based on a parametric representation (assumed shape) of the evolving temperature profile within the ice and on the integral heat budget of the ice slab. Using the integral approach, the problem of solving partial differential equations (in depth and time) is reduced to solving ordinary differential equations (in time only) for the quantities that specify the evolving temperature profile. These quantities are the ice surface temperature and the ice thickness.

In the full-fledged scheme outlined in Mironov et al. (2012), provision is made to account for the snow layer above the ice. Both snow and ice are modeled using the same basic concept, that is a parametric representation of the evolving temperature profile and the integral energy budgets of the ice and snow layers (see Mironov (2008) for a detailed discussion of the concept). In the current ICON configuration, snow over sea ice is not considered explicitly. The effect of snow is accounted for implicitly (parametrically) through the ice surface albedo with respect to solar radiation.

A prognostic sea-ice albedo parameterization is used. The sea-ice surface albedo is computed from a relaxation-type rate equation, where the equilibrium albedo and the relaxation (e-folding) time scale are computed as functions of the ice surface temperature. In order to account for the increase of the sea-ice albedo after snowfall events, the ice albedo is relaxed to the equilibrium “snow-over-ice” albedo. The equilibrium snow-over-ice albedo
is computed as function of the ice surface temperature, and the relaxation time scale is related to the snow precipitation rate.

The horizontal distribution of the ice cover, i.e., the existence of sea ice within a given ICON grid box and the ice fraction, is governed by the data assimilation scheme (cf. the treatment of lake ice). If an ICON grid box has been set ice-free during the initialization, no ice is created over the forecast period. If observational data indicate open water conditions for a given grid box but there was ice in that grid box at the end of the previous ICON run, ice is removed and the grid box is initialized as ice-free. The new ice is formed instantaneously if the data assimilation scheme indicates that there is sea ice in a given grid box, but there was no ice in that grid box in the previous model run. The newly formed ice has the surface temperature equal to the salt-water freezing point. The thickness of newly formed ice is computed as function of the ice fraction.

ICON utilizes a tile approach to compute surface fluxes of momentum and scalars. For the “sea-water type” grid boxes, the grid-box mean fluxes are computed as a weighted mean of fluxes over ice and over open water, using fractional ice cover \( f_i \) and fractional open-water cover \( 1 - f_i \) as the respective weights. Sea ice in a given ICON grid box is only considered if \( f_i \) exceeds its minimum value of 0.015, otherwise the grid box is treated as ice free (see parameter \( \text{frsi\_min} \) hard-coded in \text{mo\_seaice\_nwp\_f90}). Likewise, the open-water fraction less than \( \text{frsi\_min} \) are ignored, and the grid box in question is treated as fully ice-covered (\( f_i \) is reset to 1). The ice fraction is determined during the model initialization and is kept constant over the entire forecast period. If, however, sea ice melts away during the forecast, \( f_i \) is set to zero and the grid box is treated as an open-water water grid box for the rest of the forecast period (prognostic ice thickness is limited from below by a value of 0.05 m, i.e., a thinner ice is removed). The water-surface temperature of that grid box is equal to the observed value from the analysis, or is reset to the salt-water freezing point. The latter situation is encountered when a grid box was entirely covered by ice at the beginning of the forecast, but the ice melts away during the forecast.

In order to run ICON with the sea-ice parameterization scheme switched off, the namelist logical switch \( \text{lseaice} \) should set equal to .FALSE.. This configuration has not been comprehensively tested at DWD and is not recommended.

### 3.8.10. Land-Soil Model TERRA

The soil-vegetation-atmosphere-transfer component TERRA (Schrodin and Heise, 2001, Heise et al., 2006) in the ICON model is responsible for the exchange of fluxes of heat, moisture, and momentum between land surface and atmosphere. It establishes the lower boundary-condition for the atmospheric circulation model and considers the energy and water budget at the land surface fractions of grid points. Based on a multi-layer concept for the soil, TERRA considers the following physical processes at each of the tiled land-surface columns, where an uniform soil type with physical properties is assumed:

#### Radiation

- Photosynthetically active radiation (PAR) is used for plant evapotranspiration
- Solar and thermal radiation budget is considered in the surface energy budget

**Biophysical control of evapotranspiration**

- Stomatal resistance concept controls the interchange of water between the atmosphere and the plant
- One-layer vegetation intercepts and hold precipitation and dew, which lowers water input to the soil and enhances evaporation
- Roots with root-density profile determines the amount of water available for evapotranspiration in the soil
- Bare-soil evaporation is considered for land-surface fractions without plants.

**Heat and soil-water transport**

- Implicit numerical methods are used to solve the vertical soil water transport and soil heat transfer between the non-equidistant layers.
- In the operational model version seven layers are used in the soil.
- The lower boundary condition for the heat conduction equation is provided by the climatological mean temperature.
- Surface and sub-surface runoff of water is considered.
- The lower boundary condition is given by a free-drainage formulation.
- A rise of groundwater into the simulated soil column is not represented.
- Soil heat conductivity depends on soil-water content (Schulz et al., 2016).
- Freezing of soil water and melting of soil ice is considered in hydraulic active soil layers.

**Snow**

- TERRA offers a one-layer snow model (operational in ICON-NWP) and a multi-layer snow model option (for experiments).
- A prognostic snow density, and snow melting process as well as the time dependent snow albedo are considered
- Surface fractions partly covered with snow are divided in snow-free and snow-covered parts (snow tiles)

**Coupling to the atmosphere**

- Application of the turbulence scheme at the lower model boundary
- Roughness length for scalars implicitly considered by calculation of an additional transport resistance throughout the turbulent and laminar roughness layer.

TERRA requires a number of external parameter fields, see Section 2.4 for details.
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Figure 3.10: Tile approach for a grid cell containing various surface types. Patches of the same surface type within a grid box are regrouped into homogeneous classes (tiles) for which the soil and surface parameterizations are run separately.

The Tile Approach

The tile approach addresses the problem of calculating proper cell-averaged surface fluxes in the case of large subgrid variations in surface characteristics. The basic idea following Avissar and Pielke (1989) is depicted in Figure 3.10. If patches of the same surface type occur within a grid box, they are regrouped into homogeneous classes (tiles). The surface energy balance and soil physics are then computed separately for each tile, using parameters which are characteristic of each surface type (roughness length, leaf area index, albedo, ...). The atmospheric fields which enter the computations, however, are assumed uniform over the grid cell, i.e. the so-called blending height is located at the lowermost atmospheric model level. The contributions from different tiles are then areally weighted to provide the cell-averaged atmospheric forcing. Note that in this approach the geographical distribution of subgrid heterogeneities is not taken into account.

In ICON, the number of surface tiles is specified by the parameter `ntiles` (lnd_nml). Setting `ntiles=1` means that the tile approach is switched off, i.e. only the dominant land-surface type in a grid cell is taken into account. Setting `ntiles` to a value \( n > 1 \), up to \( n \) dominant land tiles are considered per grid cell. Note, however, that for \( n > 1 \) the total number of tiles \( n_{\text{tot}} \) is implicitly changed to \( n_{\text{tot}} = n + 3 \), with three additional "water" tiles classified as "open water" \( (n + 1) \), "lake" \( (n + 2) \), and "sea-ice" \( (n + 3) \). Additional snow-tiles can be switched on by setting `lsnowtile=TRUE`. In that case the total number of tiles is further expanded to \( n_{\text{tot}} = 2 \cdot n + 3 \), with the first \( n \) tiles denoting the land tiles, the second \( n \) tiles denoting the corresponding snow tiles and 3 water tiles as before.
Figure 3.11.: Tile generation for ntiles=3, for the case of a heterogeneous land surface. Outer circles show the fractional areas covered by the respective surface-type for a given grid cell. Inner circles show the selected tiles. Please note the re-scaling of fractional areas in the inner circle.

The process of tile generation in ICON works as follows: During the setup phase, all land-surface types within a grid box are ranked according to the fractional area $f$ they cover (see Figure 3.11, outer ring). For efficiency reasons, only the ntiles (typically about 3) dominating ones are represented by tiles, with the others being discarded (inner ring). If a grid cell contains non-negligible water bodies ($f > 5\%$), up to 3 more tiles are created (i.e. open water, lake, and sea-ice) even if they are not among the dominating ones. By this approach, the surface types represented by tiles can differ from grid cell to grid cell such that the full spectrum of surface types provided by the land cover data set is retained.

If the model is initialized from horizontally interpolated initial data and ntiles $> 1$, a tile coldstart becomes necessary. This can be done by setting $l_{tile\_init}=\text{.TRUE.}$ and $l_{tile\_coldstart}=\text{.TRUE.}$ in the namelist initicon.nml. Each tile is then initialized with the same cell averaged value. Note that $l_{tile\_init}=\text{.TRUE.}$ is only necessary, if the initial data come from a model run without tiles.

**Important note:**
Naive horizontal interpolation of tile-based variables is incorrect, since the dominant tiles and/or their internal ranking will most likely differ between source and target cell. Only aggregated fields can be interpolated!

3.8.11. Reduced Model Top for Moist Physics

A notable means for improving the efficiency of ICON is depicted in Figure 3.12. The switch
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**Figure 3.12.** Moist physics are switched off above `htop_moist_proc`, while tracer substepping is switched on above `hbot_qvsubstep`. (Remark: `hbot_qvsubstep` is allowed to be lower than `htop_moist_proc`)

`htop_moist_proc` (namelist `nonhydrostatic_nml`, floating-point value)

allows to switch off moist physics completely above a certain height. Moist physics include saturation adjustment, grid scale microphysics, convection, cloud cover diagnostic, as well as the transport of all water species but moisture $q_v$. Of course, moist processes should only be switched off well above the tropopause. The default setting is `htop_moist_proc=22500 m`.

One variant of the implemented horizontal transport scheme for passive scalars is capable of performing internal substepping. This means that the transport time step $\Delta t$ is split into $n$ (usually 2 or 3) substeps during flux computation. This proves necessary in regions where the horizontal wind speed exceeds a value of about 80 m s$^{-1}$. In real case applications, this mostly happens in the stratosphere and mesosphere. The recommendation for $\Delta t$ given in Section 3.7.1 then exceeds the numerical stability range of the horizontal transport scheme. To stabilize the integration without the need to reduce the time step globally, transport schemes with and without internal substepping can be combined. The switch

`hbot_qvsubstep` (namelist `nonhydrostatic_nml`, floating-point value)

indicates the height above which the transport scheme switches from its default version to a version with internal substepping. The default value is `hbot_qvsubstep=22500 m`.

Note that substepping is only performed for a particular tracer if a suitable horizontal transport scheme is chosen. The horizontal transport scheme can be selected individually for each tracer via the namelist switch `ihadv_tracer (transport_nml)`. Variants of the transport scheme with internal substepping are indicated by a two-digit number (i.e. 22, 32, 42, 52). These variants mostly differ w.r.t. the accuracy of the polynomial reconstruction used for the flux estimation. A linear reconstruction is used by the variant 22, whereas 52 uses a cubic reconstruction. See Section 3.6.5 for additional details regarding the transport algorithm.
If moist physics are switched off above 22.5 km (default for NWP applications), internal substepping only needs to be applied for specific humidity $q_v$, since the advection of all other moisture fields is switched off anyway. However, be aware that you must explicitly enable internal substepping if moisture physics are not switched off, or if other (non-microphysical) tracers are added to the simulation (see, e.g., Chapter 10).

### 3.9. Variable Resolution Modeling

ICON has the capability for static mesh refinement in horizontal directions. This is realized through a *multi-grid approach* which means that one or more additional high resolution (child) domains can be overlaid on coarser (parent) domains. In the following we will make frequent use of the notion *parent* and *child*, in order to illustrate the relationship amongst multiple domains.

The grid spacing factor between the parent domain and the child domain is fixed to a factor of 2, i.e. each parent triangle is split into 4 child triangles. Correspondingly, in case of nested setups, the time step $\Delta t$ is multiplied by a factor of 0.5 for each nesting level. Note that the time step $\Delta t$ needs to be specified for the base (i.e. coarsest) domain only. The (hard-coded) adaption for nested regions is done automatically.

The multi-grid approach easily allows for switching domains on or off at runtime, as well as intertwining 1-way and 2-way nested domains. 2-way as opposed to 1-way nesting means that the solution on the child domain is transferred back to the coarser parent domain every time step by means of a feedback mechanism which is described below.

The multi-grid approach closely resembles traditional two-way nesting and has to be distinguished from recent *uni-grid* approaches, where in special areas of interest more cells are added to an existing grid (*h-refinement*). Atmospheric models capable of static h-refinement are e.g. CAM-SE (Zarzycki et al., 2014) and MPAS-A (Skamarock et al., 2012). The basic multi-grid example shown in Figure 3.13 contains one global domain and one regional domain over Europe. It closely resembles the operational setup currently used at DWD.
3.9.1. Parent-Child Coupling

This section describes how the exchange of information between a parent and a child domain is realized, and which information (i.e. which fields) is interchanged.

As shown in Figure 3.14, a nested domain can conceptually be split into three areas: A boundary interpolation zone (red), a nudging zone (blue) and a feedback zone (light gray). Prognostic computations are restricted to the latter two. In case of 2-way nesting, the nudging zone does not exist and the feedback zone borders on the boundary zone. While the width of the boundary interpolation zone is fixed to 4 cell rows, the width of the nudging zone can be changed with the namelist switch `nudge_zonewidth (interpol_nml).

Once the model state on the parent domain $M_p$ has been updated from time step $n$ to $n+1$, the states $M_p^n$ and $M_p^{n+1}$ are used to update the boundary zone on the child domain. By this, the necessary lateral boundary data (forcing) can be provided for integrating the model on the nested domain from state $M_c^n$ to $M_c^{n+1}$.

In the feedback zone, the updated model state of the child domain is transferred (interpolated) back to the parent domain. By this, the parent and child domain remain closely coupled, and the simulation on the parent domain can benefit from the high-resolution results of the child domain.

In the nudging zone, which is only active for 1-way nesting, upsampled prognostic fields of the child domain are nudged towards the model state of the parent domain. A similar method is applied in limited area mode (LAM). It is further explained in Section 6.2.

Further details on the boundary update and the feedback mechanism are given in the following.

![Figure 3.13: Basic example of a multi-grid setup, consisting of a global ICON domain and a child domain over Europe with half grid spacing. This is similar to the deterministic forecast setup that is operationally used at DWD with a horizontal grid spacing of 13 km globally and 6.5 km in the child domain.](image)
Boundary Update: Parent → Child

The overall task of the boundary update mechanism is to provide the child domain with up-to-date lateral boundary conditions at each child time step. Boundary conditions are required for the following set of prognostic variables: \( v_n, w, \rho, \theta_v, q_k \). In order to avoid that interpolated values of \( \rho \) enter the solution of the continuity equation in the dynamical core, an extended set of variables is used in the boundary update mechanism, namely: \( v_n, w, \rho, \theta_v, q_k, \langle F_m \rangle \). Here, \( \langle F_m \rangle = \langle \rho v_n \rangle \) denotes a temporal average of the mass flux over the dynamic substeps. Since the continuity equation is solved in flux form (see Eq. (3.4)), making use of \( \langle F_m \rangle \) implies that interpolated values of \( \rho \) are not required for prognosing \( \rho \) on the child domain.

In general, the boundary update works as follows: Let \( \psi_p^n, \psi_p^{n+1} \) denote any of the above variables on the parent domain at time step \( n \) and \( n + 1 \), respectively. Once the model state on the parent domain \( \mathcal{M}_p \) has been updated from \( n \) to \( n + 1 \), the time tendency

\[
\frac{\partial \psi_p}{\partial t} = \frac{\psi_p^{n+1} - \psi_p^n}{\Delta t_p}
\]

is diagnosed. Both, \( \psi_p^n \) and the tendency \( \frac{\partial \psi_p}{\partial t} \) are then interpolated (downscaled) from the parent grid cells/edges to the corresponding cells/edges of the child’s boundary zone. With \( \mathcal{T}_{p\rightarrow c} \) denoting the interpolation operator, we get

\[
\psi_c^n = \mathcal{T}_{p\rightarrow c} (\psi_p^n)
\]

\[
\frac{\partial \psi_c}{\partial t} = \mathcal{T}_{p\rightarrow c} \left( \frac{\partial \psi_p}{\partial t} \right)
\]

**Figure 3.14.** General structure of a nested domain. Red: boundary interpolation zone consisting 4 cell rows. Blue: nudging zone, which is only active for 1-way nesting. Light-gray: feedback zone. Prognostic computations are performed in the feedback and nudging zone.
3.9 Variable Resolution Modeling

Since the time step on the child domain $\Delta t_c$ is only half of that on the parent domain, two integration steps are necessary in order to reach the model state $M^{n+1}_c$. The interpolated tendency is used to update the lateral boundary data after the first physics step and after each dynamics substep. E.g. the lateral boundary conditions for the first and second integration step read $\psi^n_c$ and $\psi^n_c + 0.5 \Delta t_p \partial \psi / \partial t|_c$, respectively.

Note that in order to reduce interpolation errors above steep orography, for the thermodynamic variables $\rho$ and $\theta_v$ perturbations from reference values, rather than the full values are interpolated to the child domain.

Regarding the interpolation operator $I_{p \rightarrow c}$ we distinguish between cell based and edge-based variables. For cell based variables a 2D horizontal gradient is reconstructed at the parent cell center by first computing edge-normal gradients at edge midpoints, followed by a 9-point reconstruction of the 2D gradient at the cell center based on radial basis functions (RBF, Narcovich and Ward (1994)). The interpolated value at the child cell center is then calculated as

$$\psi_c = \psi_p + \nabla \psi_p \cdot d(p,c),$$

with $\nabla \psi_p$ denoting horizontal gradient at the parent cell center, and $d(p,c)$ the distance vector between the parent and child cell center. The same operator is applied to cell based tendencies.

Regarding the interpolation of edge based variables (i.e. $\partial v_n / \partial t$, $\partial F_n / \partial t$, $\langle F_m \rangle$), we distinguish between outer child edges that coincide with the edges of the parent cell, and inner child edges.

Edge-normal vector components $\phi$ at the inner child edges are reconstructed by means of a 5-point RBF reconstruction. The 5-point stencil comprises the edges of the two parent cells sharing the edge under consideration.

For the outer child edges a more elaborate reconstruction is applied, in order to assure that the mass flux across the parent edge is equal to the sum of the mass fluxes across the two child edges. To do so, we first reconstruct the vector quantity under consideration at the parent edge vertices using RBF, from which the gradient tangential to parent edge can be computed. The edge-normal vector component at the child edge is then computed as

$$\phi_c = \phi_p + \nabla_t \phi_p \cdot d(p,c),$$

with $\nabla_t \phi_p$ denoting the gradient of the edge-normal vector component tangent to the parent edge, and $d(p,c)$ the distance vector between the parent and child edge midpoints.
Since \( d(p, c_1) = -d(p, c_2) \) holds on the ICON grid, with \( c_1, c_2 \) denoting the child cell’s edge midpoints, the above mentioned mass flux consistency is ensured.

The interpolated mass fluxes valid for the current time step can be re-computed from the interpolated average mass flux \( \langle F_m \rangle_c \), and the corresponding time tendency. These are prescribed at the outermost edges which separate the boundary zone from the prognostic zone. Since the continuity equation is solved in flux form (see Eq. (3.4)), this implies that interpolated values of \( \rho \) are not required for prognosing \( \rho \) on the child domain.

**Feedback: Child \( \rightarrow \) Parent**

If two-way nesting is selected (\texttt{lfeedback=.TRUE., namelist grid nml}), the model state \( M_{p}^{n+1} \) on the parent domain is relaxed towards the updated model state \( M_{c}^{n+1} \) on the child domain every physics time step. In the following we will refer to this as \textit{relaxation-type feedback}. It is applied to the prognostic variables \( v_n, w, \theta_v, \rho \) as well as to the prognostic, non-sedimenting mass fractions \( q_v, q_c, q_i^6 \). Let \( \psi \) denote any of the above variables. Conceptually, the method can be divided into three major steps:

1. **Upscaling**: The updated field \( \psi_{c}^{n+1} \) on the child domain is interpolated (upscaled) to the parent domain.

2. **Increment computation**: The difference between the solution on the parent domain \( \psi_p^{n+1} \) and the upscaled solution \( \psi_{c \rightarrow p}^{n+1} \) is computed.

3. **Relaxation**: The solution on the parent domain is relaxed towards the solution on the child domain. The relaxation is proportional to the increment computed in step two.

By way of example, we will mathematically describe the feedback mechanism for the variables \( \rho \) and \( \rho q_k \). Other variables are handled in a very similar manner.

The feedback mechanism for \( \rho \) can be cast into the following form:

\[
\rho_p^* = \rho_p^{n+1} + \Delta t_p \left( \mathcal{I}_{c \rightarrow p}(\rho_c^{n+1}) - \rho_p^{n+1} \right) \tag{3.44}
\]

Here \( \rho_p^{n+1} \) denotes the density in the parent cell, which has already been updated by dynamics and physics. The superscript \( "*" \) indicates the final solution, which includes the increment due to feedback. \( \Delta t_p \) is the fast physics time step on the parent domain, and \( \tau_{fb} \) is a user-defined relaxation time scale which has a default value of \( \tau_{fb} = 10800 \text{ s} \). This value is motivated by the wish to exclude small scale transient features from the feedback, but to capture synoptic-scale features. The relaxation time scale can be adjusted by means of the namelist variable \texttt{fbk relax timescale (grid nml)}. Finally, \( \mathcal{I}_{c \rightarrow p}(\rho_c^{n+1}) \) denotes a horizontal interpolation operator which upscales the child domain solution to the parent domain.

---

\(^6\)Note that when programming ICON the feedback mechanism can easily be switched on for additional tracers by adding the meta-information \texttt{lfeedback=.TRUE.} to the corresponding \texttt{add_ref}-call, see also Section 8.3.
The interpolation operator is defined as

\[ I_{c \rightarrow p}(\rho_c^{n+1}) = \sum_{i=1}^{4} \alpha_i \left( \rho_{c,i}^{n+1} - \Delta \rho_{corr} \right), \]

with \( \alpha_i \) denoting the interpolation weight for the \( i \)-th child cell. It basically performs a proper weighting of those 4 child cell center values that are connected to the parent cell under consideration. Both bilinear and area-weighted interpolation methods are available, with the former being the default choice. In order to account for differences in the vertical position of the child and parent cell circumcenters, the correction term \( \Delta \rho_{corr} \) has been introduced. At locations with noticeable orography, cell circumcenter heights at parent cells can differ significantly from those at child cells. If this is not taken into account, the feedback process will introduce a non-negligible bias in the parent domain’s mass field. The correction term is given by

\[ \Delta \rho_{corr} = (1.05 - 0.005 I_{c \rightarrow p}(\theta_v^{n+1})) \Delta \rho_{ref,p}, \]

with the difference in the reference density field

\[ \Delta \rho_{ref,p} = \sum_{i=1}^{4} (\alpha_i \rho_{ref,c,i}) - \rho_{ref,p}, \]

and the upscaled perturbation value of \( \theta_v \) denoted by

\[ I_{c \rightarrow p}(\theta_v^{n+1}) = \sum_{i=1}^{4} \alpha_i \left( \theta_{v,c,i}^{n+1} - \theta_{v,ref,i} \right). \]

The term \( \Delta \rho_{ref,p} \), which is purely a function of the parent-child height difference can be regarded as a first order correction term. In order to minimize the remaining mass drift, the empirically determined factor \((1.05 - 0.005 I_{c \rightarrow p}(\theta_v^{n+1}))\) was added, which introduces an additional temperature dependency. Note that the factor 0.005 is close to near surface values of \( \frac{\partial \rho}{\partial \theta} \) which can be derived from the equation of state.

Care must be taken to ensure that the feedback process retains tracer and air mass consistency. To this end, feedback is not implemented for tracer mass fractions directly, but for partial densities. In accordance with the implementation for \( \rho \), we get

\[ (\rho q_x)^*_{p} = (\rho q_x)^{n+1}_{p} + \frac{\Delta t_p}{\tau_{fb}} \left[ I_{c \rightarrow p}(\rho q_x^{n+1}_c) - (\rho q_x)^{n+1}_{p} \right] \]

\[ \text{(3.45)} \]

with

\[ I_{c \rightarrow p}(\rho q_x^{n+1}_c) = \sum_{i=1}^{4} \alpha_i \left( \rho_{c,i}^{n+1} - \Delta \rho_{corr} \right) q_{x,c,i}. \]

Mass fractions are re-diagnosed thereafter:

\[ q_{x,p} = \frac{(\rho q_x)^*_{p}}{\rho_{p}^*} \]

When summing Eq. (3.45) over all partial densities, Eq. (3.44) for the total density is recovered. Feedback of other prognostic variables is done in a very similar manner. Note, however, that for \( \theta_v \), the upscaling is done for the perturbation from the reference state, in order to reduce numerical errors over steep mountains. In the case of \( v_n \), some numerical diffusion is added to the resulting feedback increment in order to damp small-scale noise.
3. Model Description

Figure 3.15.: Schematic of a global domain with two two-way nested domains nested into each other. The processing sequence for integrating the model from time step \(n\) to \(n+1\), is shown in the flowchart at the lower left. Red arrows indicate boundary data interpolation from parent to child, blue arrows indicate the feedback operation from child to parent and black arrows indicate the time integration on the respective domain.

3.9.2. Processing Sequence

So far, we concentrated on the information exchange between individual parent and child domains. Nothing has been said about the processing sequence if more than one nested domain, or even repeatedly nested domains are involved. Figure 3.15 provides a common example where a global domain is accompanied by two two-way nested domains nested into each other. The global domain is schematically depicted at the bottom, whereas the nested domains are vertically staggered on top of it. The two blueish regions show the boundary interpolation zone of the nests and the feedback zone. The integration time step on the global domain is \(\Delta t\), whereas the time step is reduced by a factor of 2 when moving to the next higher grid level.

The processing sequence for integrating the model from time step \(n\) to \(n+1\) is shown in the flowchart at the lower left. The various domains are ordered top to bottom. The dots indicate the model state \(M_x\) for the different domains, red and blue arrows indicate boundary data interpolation and feedback, respectively, and black arrows indicate time integration.

The basic processing sequence is as follows:

- First, a single time step with \(\Delta t\) is performed on the global domain which results in an updated model state indicated by the white circle.

- It is followed by boundary data interpolation to the first nested domain (red arrow). Then, the model is integrated on nested domain 1 over the time interval \(\Delta t/2\).
Since there exists a second nested domain within nest 1, lateral boundary fields based on the model state $M^{n+1/2}_{c1}$ are interpolated to the second nested domain. Then, the model is integrated on the nested domain 2 over two times the time interval $\Delta t/4$, resulting in the model state $M^{n+1/2}_{c2}$.

Now, feedback is performed from nested domain 2 back to nested domain 1 (blue arrow), which results in an updated model state $M^{n+1/2}_{c1}$ on nested domain 1 (gray circle). Now, the model is again moved forward in time on nested domain 1 to reach $M^{n+1}_{c1}$.

This is followed by a second lateral boundary data interpolation for the nested domain 2. Finally, nested domain 2 is integrated in time again, to reach its final state $M^{n+1}_{c2}$. As a last step, feedback is performed from nested domain 2 to nested domain 1, followed by feedback from nested domain 1 to the global domain.

An alternative way of visualizing the processing sequence is shown in Figure 3.16. Again, the different grid levels are indicated by horizontal lines with the coarsest grid at the top and the finest one at the bottom. The model state is indicated by a white circle. Every black circle indicates that the model has been integrated in time over one domain-specific time step. Red and blue arrows again denote boundary interpolation and feedback. Note that in cases where only one domain per grid level exists (Figure 3.16a), the processing sequence resembles a W-cycle known from classic multi-grid algorithms for solving elliptic PDEs.

![Figure 3.16](image.png)

Figure 3.16.: Processing sequence (bottom row) for two generic multi-domain setups (top row). Horizontal lines represent the different grid levels and white circles denote the current model state. A filled black circle, instead, indicates a time integration step. Boundary interpolation and feedback are visualized through red and blue arrows, respectively. Setup (a) is identical to the one shown in Figure 3.15. Setup (b) differs from setup (a) by an additional nested domain on grid level 2.
3.9.3. Vertical Nesting

The model top height can be chosen individually for each domain, with the constraint that the height of the parent domain must be larger or at least equal to the height of the child domain. Thereby it is e.g. possible to combine a global domain which extends into the mesosphere with one or several child domains that only extend into the lower stratosphere, see Figure 3.17.

However, it is not possible to choose different vertical level distributions for individual domains. The level distribution is always defined by the global domain. Lower model top heights in child domains are realized by simply neglecting a certain number of levels at the model top. Vertical refinement in the sense that vertical resolution is locally increased in a child domain is so far not possible. One possible workaround might be to repeat the model run with increased vertical resolution in limited area mode (see Chapter 6).

In order to reduce the top height for individual domains, the namelist parameter num_lev (run_nml), which specifies the number of vertical levels in each domain, must be adapted. In addition, vertical nesting must be enabled by setting lvert_nest=.TRUE.. See Section 3.4 for details.

If vertical nesting is activated, boundary conditions are automatically provided for all prognostic variables at the uppermost half level of the nested domain. Several measures are taken in order to avoid excessive reflections of vertically propagating sound and gravity waves from the domain’s model top. Further details, however, are beyond the scope of this tutorial.
3.10. Reduced Radiation Grid

In real case simulations, radiation is one of the most time consuming physical processes. It is therefore desirable to reduce the computational burden without degrading the results significantly. One possibility is to use a coarser horizontal grid for radiation than for dynamics.

The implementation is schematically depicted in Figure 3.18:

**Step 1.** Radiative transfer computations are usually performed every 30 minutes. Before doing so, all input fields required by the radiation scheme are upscaled to the next coarser grid level.

**Step 2.** Then the radiative transfer computations are performed and the resulting short wave transmissivities $\tau^{SW}$ and long-wave fluxes $F^{LW}$ are scaled down to the full grid.

**Step 3.** In a last step we apply empirical corrections to $\tau^{SW}$ and $F^{LW}$ in order to incorporate the high resolution information about albedo $\alpha$ and surface temperature $T_{sfc}$ again. This is especially important at land-water boundaries and the snow line, since here the gradients in albedo and surface temperature are potentially large.

The reduced radiation grid is controlled with the following namelist switches:

- `lredgrid_phys= .FALSE./.TRUE. (namelist grid_nml, logical value)`
  - If set to .TRUE., radiation is calculated on a coarser grid (i.e. one grid level higher)

- `radiation_grid_filename (namelist grid_nml, string parameter)`
  - Filename of the grid to be used for the radiation model. Must only be specified for the base domain, since for child domains the grid of the respective parent domain

**Figure 3.18:** Schematic showing how radiation is computed on a reduced (coarser) grid.
serves as radiation grid. An empty string is required, if radiation is computed on the full (non-reduced) grid.

Note that running radiation on a reduced grid is the standard setting for operational runs at DWD. Using the reduced radiation grid is also possible for the limited area mode ICON-LAM. In this case, both the computational grid and the reduced radiation grid are regional grids. Make sure to create the latter during the grid generation process by setting dom(:)%lwrite_parent = .TRUE., see Section 2.1.3. Internally, the coarse radiation grid is denoted by the domain index 0.
4. Running Idealized Test Cases

As opposed to real case runs, idealized test cases typically do not require any external parameter or analysis fields for initialization. Instead, initial conditions are computed within the ICON model itself, based on analytical functions. These are either evaluated point-wise at cell centers, edges, or vertices, or are integrated over the triangular control volume.

The ability to run idealized model setups serves as a simple means to test the correctness of particular aspects of the model, either by comparison with analytic reference solutions (if they exist), or by comparison with results from other models. Beyond that, idealized test cases may help the scientist to focus on specific atmospheric processes.

ICON provides a set of pre-implemented test cases of varying complexity and focus, ranging from pure dynamical core and transport test cases to “moist” cases, including microphysics and potentially other parameterizations. A complete list of available test cases can be found in the namelist documentation, mentioned below.

4.1. Namelist Input for the ICON Model

In general, the ICON model is controlled by a so-called parameter file which uses Fortran NAMELIST syntax. Default values are set for all parameters, so that you only have to specify values that differ from the default.

Assuming that ICON has been compiled successfully, the next step is to adapt these ICON namelists. The run scripts in this tutorial create a file `NAMELIST_NWP` which contains all user-defined namelist parameters, together with some substituted shell script variables. Discussing all available namelist switches is definitely beyond the scope of this tutorial. We will merely focus on the particular subset of namelist switches that is necessary to setup an idealized model run. A complete list of namelist switches can be found in the namelist documentation

`icon/doc/Namelist_overview.pdf`

Individual test cases can be selected and configured by namelist parameters of the namelist `nh_testcase.nml`. To run one of the implemented test cases, only a horizontal grid file has to be provided as input. A vertical grid file containing the height distribution of vertical model levels is usually not required, since the vertical grid is constructed within the ICON model itself, based on the set of namelist parameters described in Section 3.4.
4.2. Jablonowski-Williamson Baroclinic Wave Test

From the set of available idealized test cases we choose the Jablonowski-Williamson baroclinic wave test and walk through the procedure of configuring and running this test in ICON.

The Jablonowski-Williamson baroclinic wave test (Jablonowski and Williamson, 2006) has become one of the standard test cases for assessing the quality of dynamical cores. The model is initialized with a balanced initial flow field. It comprises a zonally symmetric base state with a jet in the mid-latitudes of each hemisphere and a quasi realistic temperature distribution. Overall, the conditions resemble the climatic state of a winter hemisphere. This initial state is in hydrostatic and geostrophic balance, but is highly unstable with respect to baroclinic instability mechanisms. Thus, it should remain stationary if no perturbation is imposed.

To trigger the evolution of a baroclinic wave in the northern hemisphere, the initial conditions are overlaid with a weak (and unbalanced) zonal wind perturbation. The perturbation is centered at (20°E, 40°N). In general, the baroclinic wave starts growing observably around day 4 and evolves rapidly thereafter with explosive cyclogenesis around model day 8. After day 9, the wave train breaks (see Figure 4.1). If the integration is continued, additional instabilities become more and more apparent especially near the pentagon points (see Section 2.1), which are an indication of spurious baroclinic instabilities triggered by numerical discretization errors. In general, this test has the capability to assess

- the diffusivity of a dynamical core,
- the presence of phase speed errors in the advection of poorly resolved waves,
- the strength of grid imprinting.

In Jablonowski et al. (2008) it is suggested to add a variety of passive tracers to the baroclinic wave test case, in order to investigate the general behavior of the advection algorithm. Questions that could be addressed are

- whether the advection scheme is monotone or positive-definite,
- how accurate or diffusive the advection scheme is,
- whether a constant tracer distribution is preserved (which checks for tracer-air mass consistency).

Four different tracer distributions are implemented, whose initial distributions are depicted in Figure 4.2. See Jablonowski et al. (2008) for further information on the initial distributions.

4.2.1. Main Switches for the Idealized Test Case

This section explains several namelist groups and main switches that are necessary for setting up an idealized model run. Settings for the Jablonowski-Williamson test case are given in red.
4.2 Jablonowski-Williamson Baroclinic Wave Test

**Figure 4.1.** Surface Pressure and 850 hPa Temperature at day 9 for the Jablonowski-Williamson test case on a global R2B5 grid.

**Namelist run_nml:**

ltestcase = .TRUE. *(namelist run_nml, logical value)*

This parameter must be set to .TRUE. for running idealized test cases.

iforcing = 0 *(namelist run_nml, integer value)*

Forcing of dynamics and transport by parameterized processes. If set to 0, forcing is switched off completely (pure dynamical core test case). This implies that all physical parameterizations (see nwp_phy_nml) are switched off automatically. If set to 3, dynamics are forced by NWP-specific parameterizations. Individual physical processes can be controlled via nwp_phy_nml, see also Table 3.3. In general, the setting of iforcing depends on the selected test case.

ldynamics = .TRUE. *(namelist run_nml, logical value)*

Main switch for the dynamical core. If set to .TRUE., the dynamical core is switched on and details of the dynamical core can be controlled via dynamics_nml, nonhydrostatic_nml and diffusion_nml. If set to .FALSE., the dynamical core is switched off completely. This is rarely needed, but can be useful for idealized tests of physics packages with prescribed dynamical forcing.

ltransport = .FALSE./.TRUE. *(namelist run_nml, logical value)*

Main switch for the transport of passive tracers. If set to .TRUE., transport
Figure 4.2: Initial tracer distributions which are available for the Jablonowski-Williamson test case. Tracer $q_3$ only depends on the latitudinal position, and tracer $q_4$ is constant.

is switched on and details of the transport schemes can be controlled via transport.nml (see Section 3.6.5 for additional help). If set to .FALSE., transport of passive tracers is switched off completely.

msg_level (namelist run.nml, integer value)
You may increase the model output verbosity by setting this namelist parameter to a higher value ($\leq 20$). This option can be particularly useful if the ICON model run fails and the cause of the error still does not become clear from the error message.

Namelist nh_testcase.nml:

nh_test_name = 'jabw' (namelist nh_testcase.nml, string parameter)
Main switch for selecting a test case. nh_test_name='jabw' selects the Jablonowski-Williamson baroclinic wave test case.

Namelist extpar.nml:

itopo = 0 (namelist extpar.nml, integer value)
If set to 1, the model tries to read topography data and external parameters from file. If set to 0, no input file is required for model initialization. Instead, all initial conditions are computed within the ICON model itself. Usually, itopo should be set to 0 for running idealized test cases.
4.2.2. Specifying the Computational Domain(s)

In the following we will explain how the Jablonowski-Williamson test case can be set up for a global domain only and, in a second step, for a global domain with nests.

**Namelist grid.nml:**

- **dynamics.grid.filename** *(namelist grid.nml, list of string parameters)*
  
  Here, the name(s) of the horizontal grid file(s) must be specified. For a global simulation without nests, of course, only a single filename is required. For a global simulation with multiple nests a filename must be specified for each domain. Note that each name must be enclosed by single quotation marks and that multiple names must be separated by a comma (see the example below).

- **dynamics.parent.grid.id** *(namelist grid.nml, list of integer values)*
  
  Comma-separated list of integer values. For each domain, the grid ID of its parent domain must be specified.

  Grid IDs start with 1 and correspond to the list given by the namelist parameter **dynamics.grid.filename**. The grid ID 0 indicates that a domain has no parent domain (i.e. the global domain). Therefore the parent grid file for domain *i* is given by **dynamics.grid.filename**(dynamics.parent.grid.id(*i*))).

  Additional explanation and an example are given in Section 2.1.4 (“Step 3: Subdomain Name and Parent Grid ID”).

**Examples**

**Example 1:** Settings for a global Jablonowski-Williamson test run without nest

```
  dynamics.grid.filename = 'icon.grid.0014.R02B05.G.nc'
  dynamics.parent.grid.id = 0
  num.lev = 40
```

**Example 2:** For a global Jablonowski-Williamson test run including a single nest

```
  dynamics.grid.filename = 'icon_grid.0014_R02B05_G.nc','icon_grid.0014_R02B05_N06_1.nc'
  dynamics.parent.grid.id = 0,1
  num.lev = 40,40
```

**Example 3:** Settings for a global Jablonowski-Williamson test run including two nests on the same nesting level (i.e. combination of Figure 4.3)

```
  dynamics.grid.filename = 'icon_grid.0014_R02B05_G.nc','icon_grid.0014_R02B05_N06_1.nc',
                         'icon_grid.0014_R02B05_N06_2.nc'
  dynamics.parent.grid.id = 0,1,1
  num.lev = 40,40,40
```
4. Idealized Tests

Figure 4.3: Location of available nests for the baroclinic wave test case. The perturbation triggering the baroclinic wave is centered at (20°E, 40°N) (red circle).

Some additional information on the grid file naming convention can be found in Section 2.1.

4.2.3. Integration Time Step and Simulation Length

The integration time step and simulation length are defined as follows:

**Namelist run.nml:**

- **dtime** = 720 (namelist run.nml, real value)
  
  Time step in seconds (for the top-most domain). Note that it is *not* necessary to specify a time step for each domain. For each nesting level, the time step is automatically divided by a factor of two. More details on ICON’s time step will be given in Section 3.7.1.

- **nsteps** = 1200 (namelist run.nml, integer value)
  
  Number of time steps.

Output is controlled by the namelist group output.nml. It is possible to define more than one output namelist and each output namelist has its own output file attached to it. For example, the run script that is used in Exercise C.4.1 contains three output namelists. The details of the model output specification are discussed in Section 7.1.
5. Running Real Data Test Cases

In this chapter you will learn about how to initialize and run the ICON model in a realistic NWP setup. The namelist settings to start from a DWD Analysis and from an IFS Analysis are discussed.

5.1. Model Initialization

The necessary input data to perform a real data run have already been described in Chapter 2. These include

- grid files, containing the horizontal grid information,
- external parameter files, providing information about the Earth’s soil and land properties, as well as climatologies of atmospheric aerosols, and
- initial data (analysis) for atmosphere, land and sea.

ICON is capable of reading analysis data from various sources (see Section 2.2), including data sets generated by DWD’s Data Assimilation Coding Environment (DACE) and interpolated IFS data. In the following we provide some guidance on how to set up real data runs, depending on the specific data set at hand.

Note that ICON aborts during the setup phase, if any of the required input files has not been found.

Therefore, as a first step, check the filenames (and soft links) for the model input files, cf. Section 2. Also make sure that the input data and grid files match. For example, take a look at the global attributes number_of_grid_used and uuidOfHGrid of the global grid file. These values have to match the corresponding attributes of the external parameters and initial data file, see Section 2.1.5.

5.1.1. Basic Settings for Running Real Data Runs

Most of the main switches, that were used for setting up idealized test cases, are also important for setting up real data runs. As many of them have already been discussed in Section 4.1, we will concentrate on their settings for real data runs. Settings appropriate for the exercises on this subject (see Ex. C.5.1) are highlighted in red.
Specifying Model Start and End Dates (Namelist time_nml)

For real case runs it is important that the user specifies the correct start date and time of the simulation. This is done with `ini_datetime_string` using the ISO8601 format.

\[ \text{ini_datetime_string} = \text{YYYY-MM-DDThh:mm:ssZ} \]

— This must exactly match the validity time of the analysis data set!

Wrong settings lead to incorrect solar zenith angles and wrong external parameter fields. Setting the end date and time of the simulation via `end_datetime_string` is optional. If `end_datetime_string` is not set, the user has to set the number of time steps explicitly in `nsteps (run_nml)`, which is otherwise computed automatically.

In ICON there coexist two equally usable ways to control the experiment start and end date – without a compelling reason, though. These two alternatives are listed in the following.

Namelist run_nml:

<table>
<thead>
<tr>
<th>time step</th>
<th>dtime</th>
<th>modelTimeStep</th>
</tr>
</thead>
</table>

Namelists time_nml (left) and master_time_control_nml (right):

| experiment start | ini_datetime_string | experimentStartDate |
| experiment stop  | end_datetime_string | experimentStopDate  |

Please note that the data types of the above-mentioned namelist parameters differ. The parameters that are listed on the right are consistently based upon the ISO 8601 representations of dates and time spans. However, `dtime` must be specified in seconds.

General Settings (Namelist run_nml)

\[ \text{ltestcase} = \text{.FALSE.} \]

This parameter must be set to `.FALSE.` for real case runs.

\[ \text{iforcing} = 3 \]

A value of 3 means that dynamics are forced by NWP-specific parameterizations.

\[ \text{ldynamics} = \text{.TRUE.} \]

The dynamical core must, of course, be switched on.

\[ \text{ltransport} = \text{.TRUE.} \]

Tracer transport must be switched on. This is necessary for the transport of cloud and precipitation variables. Details of the transport schemes can be controlled via the namelist transport_nml (see Section 3.6.5).
5.1 Model Initialization

Specifying the Horizontal Grid (Namelist grid_nml)

dynamics_grid_filename (namelist grid_nml, list of string parameters)
Here, the name(s) of the horizontal grid file(s) must be specified. For a global simulation without nests, of course, only a single filename is required. For a global simulation with multiple nests, a filename must be specified for each domain. Note that each name must be enclosed by single quotation marks and that multiple names must be separated by a comma (see Section 4.2.2 and the examples therein).

dynamics_parent_grid_id (namelist grid_nml, list of string parameters)
Comma-separated list of integer values. For each domain, the grid ID of its parent domain must be specified.

Grid IDs start with 1 and correspond to the list given by the namelist parameter dynamics_grid_filename. Additional explanation and an example are given in Section 2.1.4 (“Step 3: Sub-domain Name and Parent Grid ID”). If a domain has no parent domain this is indicated by the value 0 (e.g. the global domain).

radiation_grid_filename (namelist grid_nml, string parameter)
If the radiative transfer computation should be conducted on a coarser grid than the dynamics (one level coarser, effective mesh size $2\Delta x$), the name(s) of the grid file(s) to be used for radiation must be specified here. See Section 3.10 for further details.

Specifying External Parameters (Namelist extpar_nml)

itopo= 1 (namelist extpar_nml, integer value)
For real data runs this parameter must be set to 1. The model now expects one file per domain from which it tries to read topography data and external parameters.

extpar_filename (namelist extpar_nml, string parameter)
Filename(s) of input file(s) for external parameters. If the user does not provide namelist settings for extpar_filename, ICON expects one file per domain to be present in the experiment directory, following the naming convention

```
extpar_filename = "extpar_<gridfile>.nc"
```

The keyword <gridfile> is automatically replaced by ICON with the grid filename specified for the given domain (dynamics_grid_filename). As opposed to the grid-file specification namelist variables (see above), it is not allowed to provide a comma-separated list. Instead, the usage of keywords provides full flexibility for defining the filename structure.

By changing the above setting, the user has full flexibility with respect to the filename structure. The following keywords are allowed for the namelist parameter extpar_filename:
5. Real-Data Tests

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| <path> | model base directory | (namelist parameter model_base_dir, namelist master_nml) |
| <gridfile> | grid filename for the given domain | (dynamics_grid_filename) |
| <nroot> | grid root division $R$ | (single digit) |
| <nroot0> | grid root division $R_{xx}$ | (two digits) |
| <jlev> | grid bisection level $B_{yy}$ | (two digits) |
| <idom> | domain number | (two digits). |

Specifying the Initialization Mode (Namelist initicon.nml)

ICON provides different real data initialization modes which differ in terms of the expected input fields and number of input files. Thereby ICON is able to handle analysis products from different models. The mode in use is controlled via the namelist switch init_mode.

init_mode (namelist initicon.nml, integer value)

It is possible to

- start from (interpolated) uninitialized DWD analysis without the IAU procedure: init_mode = 1
- start from interpolated IFS analysis: init_mode = 2
- start atmosphere from interpolated IFS analysis and soil/surface from interpolated ICON/GME fields: init_mode = 3
- start from non-interpolated, uninitialized DWD analysis, and make use of the IAU procedure to filter initial noise: init_mode = 5
- start from interpolated initialized ICON analysis with subsequent vertical remapping: init_mode=7

The most relevant modes are mode 1, 2, 5 and 7. They will be explained in more detail below.

ICON supports NetCDF and GRIB2 as input format for input fields. In this context it is important to note that the field names that are used in the input files do not necessarily coincide with the field names that are internally used by the ICON model. To address this problem, an additional input text file is provided, a so-called dictionary file. This file translates between the ICON variable names and the corresponding GRIB2/NetCDF short names.

Generally the dictionary is provided via the following namelist parameter:

ana_varnames_map_file (namelist initicon.nml, string parameter)

Filename of the dictionary for mapping between internal names and GRIB2/NetCDF short names. An example can be found in icon/run/ana_varnames_map_file.txt.
5.1 Model Initialization

5.1.2. Starting from Uninitialized DWD Analysis

This analysis product is rarely the optimal choice for model initialization, as it generates a significant amount of spurious noise during the first few hours of a model run (see Figure 2.6). Nevertheless, this mode is described for completeness. The process of obtaining the uninitialized DWD analysis for non-incremental update and its content is described in Section 2.2.1 (Option 2).

Model initialization is basically controlled by the following three namelist parameters:

\texttt{init\_mode = 1 (namelist initicon\_nml, integer value)}

To start from uninitialized DWD analysis data (without incremental analysis update), the initialization mode must be set to 1.

In that case the ICON model expects two input files per domain. One containing the ICON first guess (3 h forecast) fields, which served as background fields for the assimilation process. The other contains the analysis fields produced by the assimilation process. See Table 2.2 for a list of variables.

\texttt{dwdfg\_filename (namelist initicon\_nml, string parameter)}

Filename of the DWD first guess input file.

\texttt{dwdana\_filename (namelist initicon\_nml, string parameter)}

Filename of the DWD analysis input file.

Remember to make sure that the validity date for the first guess and analysis input file is the same and matches the model start date given by \texttt{ini\_datetime\_string}.

Input filenames need to be specified unambiguously, of course. By default, if the user does not provide namelist settings for \texttt{dwdfg\_filename} and \texttt{dwdana\_filename}, the filenames have the form

\texttt{dwdfg\_filename = "dwdFG\_R<nroot>B<jlev>_DOM<idom>\_nc"}
\texttt{dwdana\_filename = "dwdana\_R<nroot>B<jlev>_DOM<idom>\_nc"}

This means, e.g., that the first guess filename begins with “dwdFG”, supplemented by the grid spacing $RxB_yy$ and the domain number $DOM_{ii}$. Filenames are treated case sensitively.\footnote{More precisely this behavior depends on the file system: UNIX-like file systems are case sensitive, but the HFS+ Mac file system (usually) is not.}

By changing the above setting, the user has full flexibility with respect to the filename structure. The following keywords are allowed for the namelist parameters \texttt{dwdfg\_filename}, \texttt{dwdana\_filename} and \texttt{ifs2icon\_filename} (for the latter see Section 5.1.5):

\begin{itemize}
  \item \texttt{<path>} model base directory
  \item \texttt{<nroot>} grid root division $R_x$ (single digit)
  \item \texttt{<nroot0>} grid root division $R_{xx}$ (two digits)
  \item \texttt{<jlev>} grid bisection level $B_yy$ (two digits)
  \item \texttt{<idom>} domain number (two digits).
\end{itemize}
5.1.3. Starting from Uninitialized DWD Analysis with IAU

As described in Section 2.2.1, IAU is a means to reduce the initial noise which typically results from small scale non-balanced modes in the analysis data set. Combining this analysis product with IAU is the preferred method in cases where the horizontal and vertical grid of the intended forecast run exactly match with that of the analysis. Since no horizontal interpolation is required, the forecast run can make use of the surface tile information which is specific to this analysis product. Moreover, this product exhibits the smallest noise level during model start.

The process of obtaining the uninitialized analysis for IAU and its content is described in Section 2.2.1 (Option 1).

Model initialization is basically controlled by the following namelist parameters:

\texttt{init\_mode = 5 (namelist initicon\_nml, integer value)}

To start from DWD analysis data with IAU, the initialization mode must be set to 5.

ICON again expects two input files. One containing the ICON first guess, which typically consists of a 1.5 h forecast taken from the assimilation cycle (as opposed to a 3 h forecast used for the non-IAU case). The other contains the analysis fields (mostly increments) produced by the assimilation process. See Table 2.1 for a list of variables.

\texttt{dwdfg\_filename (namelist initicon\_nml, string parameter)}

Filename(s) of the DWD first guess input file(s) for each domain. See Section 5.1.2 for an explanation of the filename structure.

\texttt{dwdana\_filename (namelist initicon\_nml, string parameter)}

Filename(s) of the DWD analysis input file(s) for each domain. See Section 5.1.2 for an explanation of the filename structure.

The behavior of the IAU procedure is controlled via the namelist switches \texttt{dt\_iau} and \texttt{dt\_shift}:

\texttt{dt\_iau = 10800 (namelist initicon\_nml, real value)}

Time interval (in s) during which the IAU procedure (i.e. dribbling of analysis increments) is performed.

\texttt{dt\_shift = -5400 (namelist initicon\_nml, real value)}

Time (in s) by which the model start is shifted ahead of the nominal model start date given by \texttt{ini\_datetime\_string}. Typically \texttt{dt\_shift} is set to \(-0.5 \times \texttt{dt\_iau}\) such that dribbling of the analysis increments is centered around \texttt{ini\_datetime\_string}.

As explained in Section 2.2.1 and depicted in Figure 5.1, you have to make sure that the first guess is shifted ahead in time by \(-0.5 \times \texttt{dt\_iau}\) w.r.t. the analysis. The model start time \texttt{ini\_datetime\_string} must match the validity time of the analysis.
5.1 Model Initialization

Figure 5.1.: Schematic illustrating typical settings for a global ICON forecast run starting from a DWD analysis with IAU at 00 UTC. IAU is performed over a 3 h time interval (\(dt_{\text{iau}}\)), with the model start being shifted ahead of the nominal start date by 1.5 h (\(dt_{\text{shift}}\)). Validity date of first guess and analysis is 22:30 UTC and 00 UTC, respectively.

5.1.4. Starting from Initialized DWD Analysis

The initialized analysis is the product of choice in cases where the horizontal and/or vertical grid of the intended model run differs from that of the analysis. Using the uninitialized analysis for IAU is prohibited in such cases, as the horizontal interpolation of tiled surface fields makes no sense. Moreover, the initialized analysis product is less cumbersome to use, as it consists of a single file per domain, only. When compared to the standard uninitialized analysis product, spurious noise is significantly reduced (see Figure 2.6).

The process of obtaining the initialized analysis and its content is described in Section 2.2.1 (Option 3).

Model initialization is basically controlled by the following namelist parameters:

\[
\text{init\_mode} = 7 \quad \text{(namelist \textit{initicon\_nml}, integer value)}
\]

To start from initialized DWD analysis data, the initialization mode must be set to 7. If the number and or heights of the vertical levels differs between the model and the analysis, the input fields are automatically remapped in the vertical during read-in.

ICON expects a single input file:
**5. Real-Data Tests**

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`dwdg_filename (namelist initicon_nml, string parameter)`

Filename(s) of the initialized DWD analysis input file(s) for each domain. Admittedly, the nomenclature “dwdg” is a bit counter intuitive. See Section 5.1.2 for an explanation of the filename structure.

Remember to make sure that the model start time given by `init_datetime_string` matches the validity date of the input file.

**5.1.5. Starting from IFS Analysis**

No filtering procedure is currently available when starting off from interpolated IFS analysis data. The model just reads in the initial data from a single file and starts the forecast.

The process of obtaining the IFS analysis and its content is described in Section 2.2.2.

`init_mode= 2 (namelist initicon_nml, integer value)`

To start from interpolated IFS analysis data, the initialization mode must be set to 2.

`ifs2icon_filename (namelist initicon_nml, string parameter)`

ICON expects a single file per domain from which interpolated IFS analysis can be read. With this parameter, the filename can be specified. Note that for this initialization mode only input data in NetCDF format are supported. Similar to the namelist parameters `dwdg_filename` and `dwdana_filename`, which have been explained above in Section 5.1.2, the filenames have the form

```
ifs2icon_filename = "ifs2icon_R<nroot>B<jlev>_DOM<idom>.nc"
```

Remember to make sure that the model start time given by `init_datetime_string` matches the validity date of the analysis input file.

**5.2. Starting or Terminating Nested Domains at Runtime**

Starting or terminating nested domains at runtime is possible by means of the namelist parameters `start_time` and `end_time` in the namelist `grid_nml`. Model calculations for the nested domain are performed if the simulation time of the parent domain is greater or equal to `start_time` and less than `end_time`.

`start_time (namelist grid_nml, list of real values)`

Comma-separated list of integer values. For each domain, the start time relative to the experiment start date can be specified in seconds. A value of 0 for the i-th domain means that it is started at experiment start date which is either defined by `init_datetime_string` or `experimentStartDate`. If Incremental Analysis Update (IAU) is used, `start_time` must be set equal to `dt_shift (initicon_nml)` (i.e. negative), in order for the nested domain to be active from the very beginning.
5.2 Starting or Terminating Nested Domains at Runtime

end_time (namelist grid_nml, list of real values)

Comma-separated list of integer values. For each domain, the end time relative to the experiment start date can be specified in seconds. I.e. a value of 3600 specified for the \textit{i}th domain means that it is terminated one hour after experiment start.

As discussed in Section 2.2, initial data files are usually required for each nested domain. With only little loss of forecast skill, this rather tedious procedure can be overcome by starting the nested domain(s) shortly after the global domain. In that case, nested domains are initialized by parent-to-child interpolation of the prognostic fields. Note, however, that surface tile information will be lost. Surface fields on the child domain are initialized with \textit{aggregated} values interpolated from the parent domain.
5. Real-Data Tests
6. Running ICON-LAM

The most important first: Running the limited area (regional) mode of ICON does not require a separate, fundamentally different executable. Instead, ICON-LAM is quite similar to the other model components discussed so far: It is easily enabled by a top-level namelist switch

\[ \text{Namelist grid.nml: } \text{limited.area = .TRUE.} \]

Other namelist settings must be added, of course, to make a proper ICON-LAM setup. This chapter explains some of the details.

**Chapter Layout.** Some of the preprocessing aspects regarding the regional mode have already been discussed in Section 2.3. Based on these prerequisites the exercises in this chapter (see Ex. C.6.1) will explain how to actually set up and run limited area simulations.

In the following, technical details on the limited area mode are provided, in particular on how to control the read-in of initial data and boundary data.

6.1. Limited Area Mode vs. Nested Setups

In Section 3.9.1 the nesting capability of ICON has been explained. Technically, the same computational grids may be used either for the limited area mode or the nested mode of ICON\(^1\). Furthermore, both ICON modes aim at simulations with finer grid spacing and smaller scales. They therefore choose a comparable set of options out of the portfolio of available physical parameterizations.

However, there exist some differences between the regional and the one-way nested mode:

- ICON-LAM is driven by externally supplied boundary data which may come from a global model or a coarser resolution LAM that has been run in advance – that’s an obvious difference! During the simulation, boundary conditions are updated at regular time intervals by reading input files. Between two lateral boundary data samples the boundary data is linearly interpolated.

- Lateral boundary updates happen (significantly) less frequently compared to one-way nesting.

- The driving model and the limited area model may run on different computer sites. Often they also differ in terms of the governing equations as well as numerical methods used.

\(^{1}\)Here, we do not take the reduced radiation grid into account, see Section 3.10. This serves to simplify the discussion at this point.
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- ICON-LAM allows for a more flexible choice of vertical levels: Nested domains may differ from the global, “driving” grid only in terms of the top level height, but vertical layers must match between the nested and the parent domain (see Section 3.9.3). In contrast to that, the limited area mode performs a vertical interpolation of its boundary data. This is the default namelist parameter setting \texttt{itype\_latbc=1} in the namelist \texttt{limarea\_nml}. The level number and the level heights may therefore be chosen independently.

- ICON-LAM allows for a more flexible choice of the horizontal resolution. While for nested setups the increase in horizontal resolution per nesting level is constrained to a factor of 2, the resolution of the limited-area domain can be freely selected. However, resolution jumps much larger than a factor of $\sim 5$ between the forcing data resolution and the target resolution and should be avoided, since it will negatively impact the forecast quality.

6.2. Nudging in the Boundary Region

In order to prevent outward-propagating waves from reflecting back into the domain, a sponge layer is implemented along the lateral boundaries. Within this sponge layer the interior flow is relaxed towards the externally specified boundary data. In addition to this lateral boundary nudging, upper boundary nudging along the model top can be switched on by choosing \texttt{nudge\_type=1} (namelist \texttt{nudging\_nml}). The default value is 0, i.e. it is switched off. Figure 6.1 schematically depicts the partitioning of the limited area domain into the lateral boundary zone, labeled (0), the adjacent lateral nudging zone (1), the upper nudging zone (2), the nudging overlap zone (3) and the “free” model atmosphere (4). In the lateral boundary zone (0), which has a fixed width of 4 cell rows, externally supplied boundary data are simply prescribed.

The mathematical implementation of the sponge layer in the nudging zones follows the work by Davies (1976, 1983). An additional “forcing” term is added to the right hand side of the prognostic equations for $v_n$, $\theta_v$, $\rho$, and $q_v$:

$$
\psi(t) = \psi^*(t) + \alpha_{nudge} \frac{\Delta t}{\Delta \tau} \left[ \psi_{bc}(t) - \psi^*(t) \right] = \delta \psi,
$$

where $\psi_{bc}$ is the externally specified value of the prognostic variable $\psi$ at time $t$, and $\alpha_{nudge}$ is a relaxation coefficient. The nudging update $\psi - \psi^*$ depends on the time step ratio $\Delta t/\Delta \tau$ (\texttt{dyn\_substeps}), but not on the time step itself, so it might be thought of as a kind of state “shift” from $\psi^*$ to $\psi$. The nudging coefficient $\alpha_{nudge}$ gradually decreases with increasing distance from the boundary and is of the form

$$
\alpha_{nudge} = \begin{cases} 
    f_0 A_0 \exp \left( -\frac{|r-r_{0\text{ncl}}|}{\rho} \right), & \text{if } |r - r_0| \leq L \text{ (region 1 in Fig. 6.1)} \\
    B_0 \left( \frac{z-z_{\text{start}}}{2z_{\text{top}}-z_{\text{start}}} \right)^2, & \text{in region 2} \\
    \max \left\{ f_0 A_0 \exp(\cdots), B_0 (\cdots)^2 \right\}, & \text{in region 3} \\
    0, & \text{in region 4}
\end{cases}
$$
6. Nudging in the Boundary Region

6.2 Nudging in the Boundary Region

Figure 6.1.: Schematic illustration of the lateral boundary zone (blue) and the lateral and upper boundary nudging zones (gray) in the limited-area mode.

with $A_0$ the maximum relaxation coefficient in the lateral nudging zone, $L$ the width of the lateral nudging zone given in units of cell rows, $\mu$ the e-folding width given in units of cell rows, $r$ the actual cell row index beginning with 1 in the outermost cell row of the boundary zone, and $r_0$ the cell row index at which the nudging zone starts (typically $\text{grf\_bdywidth\_c}$+1, see Figure 6.1). The parameters $L$, $\mu$, and $A_0$ can be specified via $\text{nudge\_zone\_width}$, $\text{nudge\_efold\_width}$, and $\text{nudge\_max\_coeff}$ in the namelist $\text{interpol\_nml}$. The nudging zone width should at least comprise 8 (better 10) cell rows in order to minimize boundary artifacts. The tuning factor $f_0$ is either 1 for $\theta_v$ and $\rho$, or 0.5 for $v_n$ and $q_v$.

$B_0$ is the maximum nudging coefficient in the upper boundary nudging zone between the model top at height $z_{\text{top}}$ ($\text{sleve\_nml: top\_height}$) and the nudging start height $z_{\text{start}}$ ($\text{nudging\_nml: nudge\_start\_height}$). The value of $B_0$ is controlled by $\text{max\_nudge\_coeff\_vn}$ ($\text{nudging\_nml}$) for the horizontal wind $v_n$, and $\text{max\_nudge\_coeff\_thermdyn}$ for $\theta_v$ and $\rho$.

Note that positive water vapor increments $\delta q = \delta q_v > 0$ are cut to zero in supersaturated regions ($q_v > 0$) in the lateral boundary nudging zone, in order to avoid an undesirable positive feedback on the growth of the amount of cloud water. In addition, water vapor is not subject to nudging in the upper boundary nudging zone. If the nudging data from the driving model contain hydrostatic variables (i.e. hydrostatic pressure), it might be more consistent to formulate the nudging in terms of the basic hydrostatic variables: pressure and temperature. This option is controlled by the namelist switch $\text{nudge\_hydro\_pres}$ ($\text{limarea\_nml}$), which applies to both lateral and upper boundary nudging. If set to .TRUE. (default), nudging increments of the hydrostatic pressure and the temperature, $\delta p$ and $\delta T$, are applied.
are computed and transformed into virtual potential temperature and density increments following the linear mapping

\[
\delta \rho = X_\rho \delta p + Y_\rho \delta T + Z_\rho \delta q_v,
\]

\[
\delta \theta_v = X_{\theta_v} \delta p + Y_{\theta_v} \delta T + Z_{\theta_v} \delta q_v,
\]

which is motivated by the total differential of the thermodynamic state equations. The factors \(X, Y\) and \(Z\) are determined by the state before the nudging (\(\psi^*\)). Note again that water vapor increments are nonzero only in the lateral boundary nudging zone.

**Important note:** Independent of the upper boundary nudging is the treatment of the vertical velocity component \(w\) along the model top: \(w\) and corresponding vertical mass fluxes are set to zero at the uppermost half level of the computational domain. Starting from the height specified by \(\text{damp height}\) (\text{nonhydrostatic.nml}), the vertical velocity is damped towards zero following the method proposed by Klemp et al. (2008).

If upper boundary nudging is switched on (\text{nudging.nml: nudge_type=1}), “lateral boundary” data (the driving data for the nudging) have to be provided for the entire limited area domain rather than the lateral boundary region, only. This mode requires the namelist parameter \text{latbc_boundary_grid}\ (\text{limarea.nml}) defining the grid file on which the lateral boundary data are defined to be empty, i.e. \text{latbc_boundary_grid}=" ".

Running the model in regional mode is quite often accompanied by choosing a lower model top height compared to global simulations. In these cases, the neglected air mass above model top can have a noticeable impact regarding the attenuation of the incoming solar irradiance and can be the source of a small but noticeable amount of downward long-wave irradiance. To account for that in a rather ad-hoc manner, an additional model layer above model top can be added by setting \text{latm_above_top=TRUE}\ (namelist \text{nwp.phy.nml}). It is used by the radiation scheme, only. The additional layer has a (hard-coded) thickness of 1.5 times the thickness of the uppermost model layer. Currently, temperature is linearly extrapolated, assuming a vertical gradient of \(-5\) K km\(^{-1}\). For ozone, aerosols and cloud fields, a simple no-gradient condition is assumed. Despite this rather ad-hoc solution, it is suggested to activate the additional layer. Please note that this option works only in combination with a reduced radiation grid.

### 6.3. Model Initialization

The necessary input data to perform a limited area run are basically identical to those required for a global run (i.e. horizontal grid(s), initial conditions, external parameter; see Section 5.1), with the exception that lateral boundary data are required in addition in order to drive the model.

Technically it is possible to combine initial- and boundary data from different sources (e.g. one might take boundary data from IFS and initial data from ICON). In general, however, it is better to use boundary and initial data from the same source.
Dependent on the available initial data, the following initialization modes can be used in limited area mode:

\texttt{init\_mode (namelist initicon\_nml, integer value)}

- \texttt{init\_mode = 2} initialize limited area run from \textbf{IFS data}. This mode has already been described in Section 5.1.5 in the context of reading IFS analysis data.

- \texttt{init\_mode = 7} initialize limited area run from \textbf{ICON data}. This mode has already been described in Section 5.1.4 in the context of reading in DWD’s initialized analysis product.

Both modes have in common that the read-in process is followed by a vertical interpolation of the input fields to the target vertical grid. Thus the target vertical grid can be chosen independent of the vertical grid on which the input is defined. Note that in case of \texttt{init\_mode = 7} the vertical interpolation requires that the field HHL (vertical half level heights) is contained in the initial data.

\textbf{Specifics of init\_mode=2}

- Only input data in NetCDF format are supported.
- A single input file per domain is expected, containing the analysis (or, more generally, the initial state). The filename must be provided for \texttt{ifs2icon\_filename} (see also Section 5.1.5).
- The required input fields are depicted in Figure 6.2.

\textbf{Atmosphere}

\begin{align*}
\{U, V\} \quad \text{or} \quad \{W, T\} \quad \text{or} \quad \{\text{LNPS}\} & \quad \text{or} \quad \{\text{GEOSP}\} \\
\text{or} \quad \{\text{PS}\} & \quad \text{or} \quad \{\text{GEOP\_ML}\}
\end{align*}

\textbf{Soil/Surface}

\texttt{SMIL1, SMIL2, SMIL3, SMIL4, STL1, STL2, STL3, STL4, LSM, CI, GEOP\_SFC, ALB\_SNOW, SST, SKT, T\_SNOW, W\_SNOW, RHO\_SNOW, W\_I}

\textbf{Figure 6.2.}: Required variable set when initializing ICON-LAM from IFS data (i.e. \texttt{init\_mode=2}). Optional fields are marked in gray. Please note that although the field is called \texttt{W} (vertical velocity) ICON expects the content of this field to be \texttt{OMEGA} (vertical wind in a pressure based coordinate system) in case of \texttt{init\_mode=2}!
Specifics of \texttt{init\_mode=7}

- A single input file per domain is expected, containing the analysis (or, more generally, the initial state). The filename must be specified with the parameter \texttt{dwd\_ffg\_filename(initicon\_nml)}.

- As we do not make use of a second input file containing explicit analysis information, it is good practice to indicate this via the following namelist parameter.

\texttt{lread\_ana (namelist initicon\_nml, logical value)}

By default, this namelist parameter is set to \texttt{.TRUE.}. If \texttt{.FALSE.}, a separate analysis file is not required. The filename of the first guess file is specified via the \texttt{dwd\_ffg\_filename} namelist option, see Section 5.1.2.

Note that in the recent ICON version \texttt{lread\_ana=.FALSE.} is set automatically for \texttt{init\_mode=7}, if it has been forgotten by the user.

- The required input fields are listed in Table 2.3. A valid option is to use DWD’s initialized analysis product for initialization. See Section 2.2.1 for ways to obtain it.

6.4. Reading Lateral Boundary Data

The read-in of lateral boundary data is fortunately less cumbersome than the read-in of initial data, as it is based on a decision tree. The user is no longer required to select a specific mode which (hopefully) fits the data at hand. Instead, ICON scans the boundary data file and, dependent on its content, ICON diagnoses additional fields so as to obtain the internally required set of variables. The decision tree is depicted in Figure 6.3. If the provided data set does not match any of the trees, an error is thrown. As a result, ICON can handle variable sets from hydrostatic models (e.g. IFS) as well as non-hydrostatic models (e.g. COSMO, ICON) without the assistance of the user.

As apparent from the decision tree, three different variable sets can be handled. See Figure 2.9 for its specific content.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure63.png}
\caption{Decision tree for the read-in of lateral boundary data.}
\end{figure}

\textbf{Important note:} Boundary data sets originating from a non-hydrostatic model with height based vertical coordinates (e.g. COSMO or ICON) must contain the field \texttt{HHL} (vertical half level heights). It is required by the vertical interpolation procedure. Note, however, that the field only needs to be present in the boundary data set whose validity date equals the model start date.

Read-in of boundary data is controlled by the following namelist parameters:

The type of lateral boundary conditions is specified by

\texttt{itype\_latbc (namelist limarea\_nml, Integer value)}

If set to 1 time-dependent boundary conditions are used. ICON then tries to read external data files at regular time intervals from a particular location specified by
Figure 6.3: Read-in of lateral boundary data. Based on this decision tree, ICON investigates the data file contents and diagnoses additional fields.
The following fields are read additionally from file: velocity fields $\text{VN}$ (or $U$, $V$) and mixing ratios $\text{QV}$, $\text{QC}$, $\text{QI}$ (optional: $\text{QR}$, $\text{QS}$). The fields $W$ and $\text{OMEGA}$ are optional; if they are unavailable, the vertical wind is initialized with zero.
For the input from a pressure based coordinate system (right branch), note that ICON expects the field $\text{OMEGA}$ under the name $W$.

latbc_filename and latbc_path (see below).
If set to 0, time-constant lateral boundary conditions are used which are derived from the initial conditions.

Boundary data is read at regular time intervals. This is specified by the following namelist parameter:
dtime_latbc (namelist limarea_nml, floating-point value)
Time difference in seconds between two consecutive boundary data sets. At intermediate times, boundary conditions are computed by linear interpolation in time.

6.4.1. Naming Scheme for Lateral Boundary Data

Naturally, the sequence of lateral boundary data files must satisfy a consistent naming scheme. It is a good idea to consider this convention already during the preprocessing steps (see Section 2.3).
6. Limited Area Mode

**Filenames:** latbc_filename, latbc_path (string parameters, limarea_nml)

By default, the filenames are expected to have the following form:

"prepiconR<nroot>B<jlev>_<y><m><d><h>.nc"

Here, several keywords are used which are further explained below. This naming scheme can be flexibly altered via the namelist parameter latbc_filename (namelist limarea_nml) using the available keywords. The absolute path to the boundary data can be specified with latbc_path (string parameter, limarea_nml).

By changing the above setting, the user has full flexibility with respect to the filename structure. The following keywords are allowed for the namelist parameter latbc_filename:

- `<nroot>`: grid root division Rz (single digit)
- `<nroot0>`: grid root division Rzz (two digits)
- `<jlev>`: grid bisection level Byy (two digits)
- `<dom>`: domain number (two digits)
- `<y>`: year (four digits)
- `<m>`: month (two digits)
- `<d>`: day in month (two digits)
- `<h>`: hour (UTC) (two digits)
- `<min>`: minutes (UTC) (two digits)
- `<sec>`: seconds (UTC) (two digits)
- `<ddhhmmss>`: elapsed days, hours, minutes and seconds since ini_datetime_string or experimentStartDate (each two digits)
- `<dddhh>`: elapsed days and hours since ini_datetime_string or experimentStartDate (three digits day, two digits hours).

**Field names:** latbc_varnames_map_file (namelist limarea_nml, string)

ICON supports NetCDF and GRIB2 as input format for boundary fields. Field names in input files do not necessarily coincide with internal ICON field names. Hence, an additional input text file (dictionary file) can be provided. This two-column file translates between the ICON variable names and the corresponding DWD GRIB2 short names or NetCDF variable names.

Specifying a valid dictionary file is currently mandatory, if pre-fetching of boundary data is selected num_prefetch_proc=1 (see below).

**Boundary grid:** latbc_boundary_grid (namelist limarea_nml, string)

As it has been explained in Section 2.3, the lateral boundary data can be defined on an auxiliary grid, which contains only the cells of the boundary zone for optimization purposes.

If this is the case for the applied boundary data, the filename of this grid file must be specified with this namelist parameter.
6.4.2. Pre-Fetching of Boundary Data (Mandatory)

*Pre-fetching* strives to avoid blocking of the computation due to reading of boundary data. The term denotes the reading of files ahead of time, i.e. the next input file will be processed simultaneously with the preceding compute steps. This avoids waiting for the I/O processes during the time consuming procedure of opening, reading and closing of the input files.

\[
\text{num\_prefetch\_proc} = 1 \quad \text{(namelist parallel\_nml, integer value)}
\]

If this namelist option is set to 1, one MPI process will run exclusively for asynchronously reading boundary data during the limited area run. This setting, i.e. the number of pre-fetching processors, can be zero or one.

Enabling the pre-fetching mode is **mandatory** for the described LAM setup.
7. Parallelization and Output

In this chapter we describe advanced settings for the namelist controlled model output.

In particular, the ICON model offers several options for internal post-processing, such as the horizontal remapping of the prognostic output onto regularly spaced (“longitude-latitude”) grids and vertical interpolation, for example on pressure levels. Another type of “output products” is ICON’s checkpointing feature which allows to restart the execution from a pre-defined point using the data stored in a file.

The chapter is concluded by an overview of the different mechanisms for parallel execution of the ICON model. These settings become important for performance scalability when increasing the model resolution and core counts.

7.1. Settings for the Model Output

Model output is enabled via the namelist run_nml with the main switch output. By setting this string parameter to the value "nml", the output files and the fields requested for output can be specified by special namelists. In the following, this procedure will be described in more detail.

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

a) The time interval between two model outputs.

b) The name of the output file.

c) The name(s) of the variable(s) to output.

d) The type of the vertical output grid (e.g. pressure levels or model levels).

e) The type of the horizontal output grid (i.e. ICON grid or geographical coordinates).

All of these parameters are set in the namelist output_nml. Multiple instances of this namelist may be specified for a single model run, where each output_nml creates a separate output file. The options d) and e) require an interpolation step. They will be discussed in more detail in Section 7.1.1.

In the following, we give a short explanation for the most important namelist parameters:

output_filename (namelist output_nml, string parameter)

This namelist parameter defines a prefix for the output filename (which may include the directory path). The domain number, level type, file number and file format extension will be appended to this prefix.
output_bounds (namelist output_nml, three floating-point values)

This namelist parameter defines the start time and the end time for the model output and the interval between two consecutive write events. The three values for this parameter are separated by commas and, by default, they are specified in seconds.

ml_varlist (namelist output_nml, character string list)

This parameter is a comma-separated list of variables or variable groups (the latter are denoted by the prefix "group:"). The ml_varlist corresponds to model levels, but all 2D variables (for example surface variables) are specified in the ml_varlist as well. It is important to note that the variable names follow an ICON-internal nomenclature. The temperature field, for example, is denoted by the character string “temp”. A list of available output fields is provided in Appendix B.

Users can also specify the variable names in a different naming scheme, for example “T” instead of “temp”. To this end, a translation table (a two-column ASCII file) can be provided via the parameter output_nml_dict in the namelist io_nml. An example for such a dictionary file can be found in the source code directory: run/dict.output.dwd.

m_levels (namelist output_nml, character string)

This character string specifies a list of model levels for which the variables and groups should be written to output. Level ordering does not matter. Allowed is a comma- (or semicolon-) separated list of integers, and of integer ranges like “10...20”. 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 = "1,3,5...10,20...(nlev-2)"

dom (namelist output_nml, integer values, comma-sep.)

Related to setups with nests, i.e. multiple domains: This namelist parameter sets the domains for which this namelist is used. If not specified (or specified as -1), this namelist will be used for all domains.

remap (namelist output_nml, integer value: 0/1)

This namelist parameter is related to the horizontal interpolation of the output to regular grids, see Section 7.1.1.

filetype (namelist output_nml, integer value: 2/4)

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. Here, the value filetype=2 denotes the GRIB2 output, while the value filetype=4 denotes the NetCDF file format.

The namelist parameter output_filename provides only partial control over the resulting filename. Complete control over the resulting filename can be achieved with the namelist parameter filename_format (namelist output_nml, character string). By default, filename_format is set to:
7. Parallelization & I/O

7.1 Settings for the Model Output

The following keywords are allowed for filename_format (non-exhaustive list):

- `<path>` model base directory
  (namelist parameter `model_base_dir`, namelist `master_nml`)
- `<physdom>` domain number (two digits)
- `<levtype>` Level type (ML, PL, HL, IL)
- `<datetime>` Date and time using the ISO8601 format.
- `<ddhhmmss>` elapsed days, hours, minutes and seconds since `ini_datetime_string` or `experimentStartDate` (each two digits)
- `<jfile>` Consecutive file id.

As it has been stated before, each `output_nml` creates a separate output file. To be more precise, there are a couple of exceptions to this rule. First, multiple time steps can be stored in a single output file, but they may also be split up over a sequence of files (with a corresponding index in the filename), cf. the namelist parameter `steps_per_file`. Second, an instance of `output_nml` may also create more than one output file if grid nests have been enabled in the model run together with the global model grid, cf. the namelist parameter `dom`. In this case, each of the specified model domains is written to a separate output file. Finally, model output is often written on different vertical axes, e.g. on model levels and on pressure levels. The specification of this output then differs only in the settings for the vertical interpolation. Therefore it is often convenient to specify the vertical interpolation in the same `output_nml` as the model level output, which again leads to multiple output files.

7.1.1. Output on Regular Grids and Vertical Interpolation

Many diagnostic tools, e.g. to create contour maps and surface plots, require a regularly spaced distribution of the data points. Therefore, the ICON model has a built-in output module for the interpolation of model data from the triangular mesh onto a regular longitude-latitude grid. Further information on the interpolation methods can be found in the database documentation (cf. Section 0.3). Furthermore, the model output can be written on a different vertical axis, e.g. on pressure levels, height levels or isentropes. In the following we will describe how to specify these options.

All of these parameters are set in the namelist `output_nml`. As it was already mentioned in Section 7.1, multiple instances of this namelist may be specified for a single model run, where each `output_nml` creates a separate output file.

The relevant namelist parameters for the `horizontal` interpolation of the output fields are:

`remap` (namelist `output_nml`, integer value 0/1)

Set this namelist parameter to the value 1 to enable horizontal interpolation onto a regular grid. This option needs to be defined combination with `reg_lat_def` / `reg_lon_def`. 

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reg_lat_def / reg_lon_def (namelist output_nml)
   Latitudes and longitudes for the regular grid points are each specified by three values:
   start, increment, end value; given in degrees. Alternatively, the user may set the
   number of grid points instead of an increment.

The relevant namelist parameters for the vertical interpolation of the output fields are:

hl_varlist / pl_varlist / il_varlist (character string lists)
   Similar to the namelist parameter ml_varlist mentioned above, these parameters
   are comma-separated lists of variables or variable groups. While the hl_varlist
   sets the output for height levels, pl_varlist defines variables on pressure levels and
   il_varlist specifies output on isentropic levels.

h_levels / p_levels / i_levels (floating point values, comma-sep.)
   Comma separated list of height, pressure, and isentropic levels for which the variables
   and groups specified in the above mentioned variable lists should be output. Height
   levels must be given in m, pressure levels in Pa and isentropes in K. Level ordering
   does not matter.

7.1.2. Output Rank Assignment

When a large number of different output files is written during the simulation, the task of
formatting and writing may put an excessive load on the output processes. The number of
output processes which share this output load can be increased by setting the num_io_procs
(parallel_nml) namelist parameter, see Section 7.3.1. If there exist multiple groups (e.g.
output files, different variable sets, output intervals, interpolation grids) then these so-
called streams will be distributed automatically over the available output processes.

stream_partitions_ml (namelist output_nml, integer)
   It may be even useful to spread the files of a single stream over multiple output
   processes. For example, when each output file is relatively large, then the subsequent
   file of this stream can be written by a different output process in order to diminish
   the risk of congestion. Please use the namelist parameter stream_partitions_ml to
   set the number of output processes among which the output files should be divided.

The distribution of output load using stream_partitions_ml is illustrated in
Fig. 7.1.

pe_placement_ml (namelist output_nml, integer array)
   This array is related to the namelist parameters num_io_procs and
   stream_partitions_ml and allows for an even more fine-tuned distribution of
   the output workload. At most stream_partitions_ml different ranks can be speci-
   fied, ranging between 0 ... (num_io_procs - 1). This explicitly assigns the output
   streams to specific PEs and facilitates a load balancing with respect to small and
   large output files.
7.2 Checkpointing and Restart

There are many reasons why a simulation execution may be interrupted prematurely or unexpectedly. The checkpoint/restart option can save you from having to start the ICON model over from the beginning if it does not finish as expected. It allows you to restart the execution from a pre-defined point using the data stored in a checkpoint file.

Activating the restart. The checkpoint/restart functionality is controlled by the following namelist parameters:

\texttt{dt\_checkpoint} (\texttt{namelist io\_nml}, floating-point value)

This parameter specifies the time interval for writing restart files. The restart files are written in NetCDF format, and their names are specified by the namelist parameter \texttt{restart\_filename}, see below.

Note that if the value of \texttt{dt\_checkpoint} resulting from the model default or user's specification is larger than \texttt{dt\_restart} (see below), then it will be automatically reset to \texttt{dt\_restart}, s.t. at least one restart file is generated during the restart cycle.

\texttt{lrestart} (\texttt{namelist master\_nml}, logical value)

If this namelist parameter is set to .\texttt{TRUE.} then the current experiment is resumed from a restart file.

Instead of searching for a specific data filename, the model reads its restart data always from a file with name \texttt{restart\_atm\_DOM01.nc} (analogously for nested domains). It is implicitly assumed that this file contains the newest restart data, because during the writing of the checkpoints this file is automatically created as a symbolic link to the latest checkpoint file.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{distribution.png}
\caption{Schematic illustrating the distribution of output load onto three output processes, using \texttt{num\_io\_procs=3} and \texttt{stream\_partitions\_ml=3}.}
\end{figure}
**restart_filename (namelist run.nml, string parameter)**

This namelist parameter defines the name(s) of the checkpointing file(s). By default, the checkpoint files (not the symbolic link) have the form

```
gridfile_restart_atm_restarttime.nc
```

**dt_restart (namelist time.nml, floating-point value)**

This parameter is in some ways related to the dt_checkpoint parameter: It specifies the length of a restart cycle in seconds, i.e. it 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.

Similar to the asynchronous output module, the ICON model (cf. Section 7.3) also offers the option to reserve a dedicated MPI task for writing checkpoint files. This feature can be enabled by setting the parameter num_restart_procs in the namelist parallel.nml to an integer value larger than 0.

**Restart modes.** Different restart write modes are available, which allow for a distributed writing and read-in of restart files, depending on the parallel setup. These different restart modes are controlled via the namelist parameter restart_write_mode (io.nml).

Allowed settings for restart_write_mode (character strings!) are:

"joint procs multifile"

All worker processes write restart files to a dedicated directory. Therefore, the directory itself represents the restart data set. The information is stored in a way that it can be read back into the model independent from the processor count and the domain decomposition.

*Read-in:* All worker processes read the data in parallel.

"dedicated procs multifile"

In this case, all the restart data is first transferred to memory buffers in dedicated restart writer processes. After that, the work processes carry on with their work immediately, while the restart writers perform the actual restart writing asynchronously. Restart processes can parallelize over patches and horizontal indices.

*Read-in:* All worker processes are available to read the data in parallel (though this is usually limited by the number of restart files).

"sync"

'Old' synchronous mode. Process # 0 reads and writes restart files. All other processes have to wait.

"async"

'Old' mode for asynchronous restart writing: Dedicated processes (num_restart_proc > 0) write restart files while the simulation continues. Restart processes can only parallelize over different patches.

*Read-in:* Processes # 0 reads while other processes have to wait.
7.3 Parallelization and Performance Aspects

As mentioned in Section 1.2.1, ICON can use two different mechanisms for parallel execution:

a) **OpenMP** – Multiple threads are run in a single process and share the memory of a single machine.
   An implementation of OpenMP ships with your Fortran compiler. OpenMP-parallel execution therefore does not require the installation of additional libraries.

b) **MPI** – Multiple ICON processes (*processing elements*, PEs) are started simultaneously and communicate by passing messages over the network. Each process is assigned a part of the grid to process.

These mechanisms are not mutually exclusive. A *hybrid* approach is also possible: Multiple ICON processes are started, each of which starts multiple threads. The processes communicate using MPI. The threads communicate using OpenMP.

7.3.1. Settings for Parallel Execution

Several settings must be adjusted to control the parallel execution:

**Namelist parallel_nml**
First, we focus on some namelist settings for the distributed-memory MPI run.

Processors are divided into

- **Worker PEs** – this is the majority of MPI tasks, doing the actual work
- **I/O PEs** – dedicated output server tasks\(^1\)
- **Restart PEs** – for asynchronous restart writing (see Section 7.2)
- **Prefetch PE** – for asynchronous read-in of boundary data in limited area mode (see Section 6.4.2)
- **Test PE** – MPI task for verification of MPI parallelization (debug option)

The configuration settings are defined in the namelist `parallel_nml`. To specify the number of output processes, set the namelist parameter `num_io_procs` to a value larger than 0, which reserves a number of processors for output. While writing, the remaining processors continuously carry out calculations. Conversely, setting this option to 0 forces the worker PEs to wait until output is finished. For the writing of the restart checkpoints (see Section 7.2), there exists a corresponding namelist parameter `num_restart_procs`.

---

\(^1\)The notation “I/O” is justified by historical arguments. In the current version of ICON, these MPI processes exclusively operate as output servers.
During start-up, the model prints out a summary of the processor partitioning. This is often helpful to identify performance bottlenecks. First of all, the model log output contains a one-line status message:

**Number of procs for**


Afterwards, the sizes of grid partitions for each MPI process are summarized as follows:

- **Number of compute PEs used for this grid:** 118
- **# prognostic cells:** max/min/avg xxx xxx xxx

Given the case that the partitioning process would fail, these (and the subsequently printed) values would be grossly out of balance.

### Batch queuing system

Apart from the namelist settings, the user has to specify the computational resources that are requested from the compute cluster. In addition to the number of MPI tasks and OpenMP threads, here the user has to set the number of cluster-connected nodes.

Increasing the number of nodes allows to use more computational resources, since a single compute node comprises only a limited number of PEs and OpenMP threads. On the other hand, off-node communication is usually more expensive in terms of runtime performance.

When using the `qsub` command to submit a script file, the queuing system PBSPro allows for specification of options at the beginning of the file prefaced by the `#PBS` delimiter followed by PBS commands (see also the comments in Appendix A). For example, to run the executable in hybrid mode on 12 nodes with 4 OpenMP threads/processes, set

```bash
#PBS -q xc_norm_h
#PBS -l select=12:ompthreads=4
#PBS -l place=scatter
#PBS -l walltime=01:00:00
#PBS -j oe
```

In more detail, the PBS keywords have the following meaning:

- `#PBS -q xc_norm_h`  
  Job for the (Haswell) compute nodes of the XC 40.
- `#PBS -l select=$NODES`  
  To specify the number of compute nodes.
- `#PBS -l place=pack`  
  If only one compute node is used.
- `#PBS -l place=scatter`  
  If more than one compute node are used.
- `#PBS -l walltime=...`  
  This directive specifies the maximum wall-clock time (real time) that a job should take.
- `#PBS -j oe`  
  Put standard error and log to the same device.
- `#PBS -N`  
  To give a special name to the job.
7.3 Parallelization and Performance Aspects

Application launch with aprun

Finally the user has to set the correct options for the application launcher, which is the `aprun` command on the Cray XC 40 platform. Here we have the following syntax:

```
aprun -n total number of MPI tasks
-N number of MPI tasks/node
-d number of threads/MPI task
-j hyperthreading (enabled=2)
-m memory/task
command
```

Best Practice for Parallel Setups

ICON employs both distributed memory parallelization and shared memory parallelization, i.e. a “hybrid parallelization”. Only the former type actually performs a decomposition of the domain data, using the de-facto standard MPI. The shared memory parallelization, on the other hand, uses OpenMP directives in the source code. In fact, nearly all `DO` loops that iterate over grid cells are preceded by OpenMP directives. For reasons of cache efficiency the `DO` loops over grid cells, edges, and vertices are organized in two nested loops: “`jb` loops” and “`jc` loops”\(^1\) Here the outer loop (“`jb`”) is parallelized with OpenMP.

There is no straight-forward way to determine the optimal hybrid setup, except for the extreme cases: If only a single node is used, then the global memory facilitates a pure OpenMP parallelization. Usually, this setup is only feasible for very small simulations. If, on the other hand, each node constitutes a single-core system, a multi-threaded (OpenMP) run would not make much sense, since multiple threads would interfere on this single core. A pure MPI setup would be the best choice then.

In all of the other cases, the parallelization setup depends on the hardware platform and on the simulation size. In practice, 4 threads/MPI task have proven to be a good choice on x86-based systems. This should be combined with the `hyper-threading` feature, i.e. a feature of the x86 architecture where one physical core behaves like two virtual cores.

Starting from this number of threads per task the total number of MPI tasks is then chosen such that each node is used to an equal extent and the desired time-to-solution is attained – in operational runs at DWD this is \(\sim 1\) h. In general one should take care of the fact that the number of OpenMP threads evenly divides the number of cores per CPU socket, otherwise inter-socket communication might impede the performance.

Finally, there is one special case: If an ICON run turns out to consume an extraordinarily large amount of memory (which should not be the case for a model with a decent memory scaling), then the user can resort to “investing” more OpenMP threads than it is necessary for the runtime performance. Doing so, each MPI process would have more memory at its disposal.

\(^1\)This implementation method is known as `loop tiling`, see also Section 8.3.
7.3.2. Mixed Single/ Double Precision in ICON

To speed up code parts strongly limited by memory bandwidth, an option exists to use single precision for variables that are presumed to be insensitive to computational accuracy – primarily the dynamical core and the tracer advection.

This affects most local arrays in the dynamical core routines, some local arrays in the tracer transport routines, the metrics coefficients, arrays used for storing tendencies or differenced fields (gradients, divergence etc.), reference atmosphere fields, and interpolation coefficients. Prognostic variables and intermediate variables affecting the accuracy of mass conservation are still treated in double precision.

To activate the mixed-precision option, the preprocessor flags `-D__MIXED_PRECISION` and `-D__MIXED_PRECISION_2` need to be specified in the configuration settings used for generating the Makefile. The latter flag is used for physics tendencies.

There exists a configure option which enables the above preprocessor flags:

```
./configure --enable-mixed
```

Note that interpolation to a latitude-longitude grid is not supported for single-precision variables; if you desire to output physics tendency fields on a regular grid for diagnostic purposes, do not set `-D__MIXED_PRECISION_2`.

7.3.3. Bit-Reproducibility

Bit-reproducibility refers to the feature that running the same binary multiple times should ideally result in bitwise identical results. Depending on the compiler and the compiler flags used this is not always true if the number of MPI tasks and/or OpenMP threads is changed in between. Usually compilers provide options for creating a binary that offers bit-reproducibility, however this is often payed dearly by strong performance losses. With the Cray compiler, it is however possible to generate an ICON binary offering bit-reproducibility with only little performance loss. The ICON binary used in this workshop gives bit-reproducible results (will be checked in Exercise C.5.7).

Bit-reproducibility is generally an indispensable feature for debugging. It is helpful

- for checking the MPI/OpenMP parallelization of the code. If the ICON code does not give bit-identical results when running the same configuration multiple times, this is a strong hint for an OpenMP race condition. If the results change only when changing the processor configuration, this is a hint for a MPI parallelization bug.
- for checking the correctness of new code that is supposed not to change the results.

7.3.4. Basic Performance Measurement

The ICON code contains internal routines for performance logging for different parts (setup, physics, dynamics, I/O) of the code. These may help to identify performance bottlenecks. ICON performance logging provides timers via the two namelist parameters `ltimer` and `timers_level` (namelist `run_nml`).

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Note for advanced users: The built-in timer output is rather non-intrusive. It is therefore advisable to have it enabled also in operational runs.

With the following settings in the namelist `run.nml`,

```plaintext
ltimer = .TRUE.
timers_level = 10
```

the user gets a sufficiently detailed output of wall clock measurements for different parts of the code:

<table>
<thead>
<tr>
<th>name</th>
<th># calls</th>
<th>total min (s)</th>
<th>total max (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>total</td>
<td>237</td>
<td>903.085</td>
<td>903.089</td>
</tr>
<tr>
<td>L integrate_nh</td>
<td>170640</td>
<td>884.128</td>
<td>892.143</td>
</tr>
<tr>
<td>L nh_solve</td>
<td>5972400</td>
<td>401.055</td>
<td>428.694</td>
</tr>
<tr>
<td>L nh_solve.veltend</td>
<td>7166880</td>
<td>36.469</td>
<td>51.376</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>physics</td>
<td>49678</td>
<td>103.107</td>
<td>104.759</td>
</tr>
<tr>
<td>L nwp_radiation</td>
<td>10030</td>
<td>40.402</td>
<td>42.985</td>
</tr>
<tr>
<td>L radiation</td>
<td>220674</td>
<td>31.845</td>
<td>34.963</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>model_init</td>
<td>711</td>
<td>59.875</td>
<td>59.876</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note that some of the internal performance timers are nested, e.g. the timer log for `radiation` is contained in `physics`, indicated by the “L” symbol. For correct interpretation of the timing output and computation of partial sums one has to take this hierarchy into account.

- The column "total max (s)" contains the maximum timing in seconds (maximum over all MPI tasks, OpenMP master thread).
- The row "model_init" contains the measurements for the model setup (allocation, read-in, etc.).
- The row "total" contains the model run-time, excluding the initialization and finalization phase.
8. Programming ICON

Just because something doesn’t do what you planned it to do doesn’t mean it’s useless.

Thomas Edison

The previous chapters’ topics have been guided by questions of how to run ICON simulations in various settings and how to control and understand the model’s characteristics. In this short chapter, instead, we will introduce ICON’s inner workings, i.e. the code layout and the most important data structures.

The description is detailed enough to make it relatively easy for the reader to modify the code. We exemplify this in Section 8.3 by implementing an own simple diagnostic.

8.1. Representation of 2D and 3D Fields

We begin with a suitable representation of two- and three-dimensional fields. Here, we refer to a discrete variable as a 2D field, if it depends on the geographical position only. A 3D field, in addition, contains a vertical dimension, associated with the grid column.

Indexing. Recalling the unstructured nature of ICON’s computational grids (see Section 2.1) there is no obvious order of the cells in a 2D array like indexing them according to longitudes and latitudes. Instead, we just order the cells in a deliberate way and index them in this order with ascending integer numbers. This means that our 2D field becomes a 1D array, referenced by the cell indices as subscript values.

Most arrays are associated with the centers of the triangular grid cells, but we do that in a similar way for the edges and vertices of the triangles. An extension to 3D fields, i.e. including a vertical dimension, results in 2D arrays, the first index being the cell (or edge or vertex) index, the second index being the height level.

Blocking. For reasons of cache efficiency nearly all DO loops over grid cells, edges, and vertices are organized in two nested loops: “jb loops” and “jc loops”. Often, the outer loop (“jb”) is parallelized with OpenMP.

With respect to the data layout, this means that the long vector is split into several chunks of a much smaller length nproma (this is a run-time parameter, defined in the namelist parallel_nml). We store the long vector in a 2D array, the first index counting
Figure 8.1.: Illustration of the 2D field representation. The original spherical domain is decomposed (light-gray: halo region). Afterwards, the long vector of grid cells is split into several chunks of a much smaller length $n_{proma}$.

the elements in a block (line index), the second index counting the blocks. The last block may be shorter since $n_{proma}$ is not necessarily a divisor of the number of cells. The blocking procedure is illustrated in the lower half of Figure 8.1. There exist auxiliary functions idx_no, blk_no and idx_1d, which help to calculate the blocked indices from the 1D array index and vice versa (declared in the module mo_parallel_config).

Finally, let us consider the 3D fields that were stored in 2D arrays, with the cell index as the first dimension and the second being the vertical coordinate. These fields will be stored in 3D arrays with the first index counting the elements in a block, the second index counting the levels and the third index counting the blocks. The reason is that the blocks are often passed one by one to some subprograms which are called in a loop over the blocks. Since Fortran stores arrays in column-major order, the data for a single $jb$ is stored contiguously in memory. Thus we can pass this chunk of data to the subprograms without any reshape of the arrays.

**Domain decomposition.** Domain decomposition is, naturally, a prerequisite for scalability on modern parallel computers. For large scale realistic ICON setups and with operational core counts in the range of tens of thousands, the use of persistent global-sized arrays is unacceptable. Each model domain is therefore distributed onto several processors\(^1\). This means that we have only certain regions of a domain on each processor.

\(^1\)In the following, we will use the generic term “Processing Element” (PE).
Each processor’s region consists of an inner portion and a lateral boundary portion. The latter may be either a lateral boundary for the entire domain or a \textit{halo region}, i.e. a lateral boundary of the partial domain which is overlaid with neighboring partial domains. The halo region (which is also known as a \textit{ghost-cell region}) is illustrated in the upper half of Figure 8.1.

The programmer is responsible for the distribution of the data among the processors and the correct communication through MPI calls. This means that all halo regions have to be updated by the neighboring partial domains. We will sketch this synchronization process in Section 8.2.4 below.

\begin{center}
\begin{tabular}{|c|}
\hline
\textbf{Keep in mind that it is the width of the halo region which defines a size limit for your stencil calculations: It is not possible to include cells in a stencil which extend beyond the halo region!} \\
\hline
\end{tabular}
\end{center}

**Index ordering.** After the domain decomposition, which takes place in the model initialization phase, each PE performs a sorting of its locally allocated cells (and edges and vertices). This \textit{local} index ordering is determined by the \texttt{refin.\texttt{xxx} ctrl} index which counts the distance from the lateral boundary in units of cell/edge/vertex rows. In particular, note the \texttt{refin.c ctrl} array which already played a role for the preparation of lateral boundary input data, see Section 2.3. Portions of the triangular cells correspond to different values of \texttt{refin.c ctrl}: the cell rows at the lateral boundary, the nudging zone, the inner cells, and the halo region. For a global domain only the two latter categories exist.

The lower part of Figure 8.2 visualizes the ordering of the grid portions in the index vector. It can be seen that the leftmost indices (i.e. the smallest subscripts) correspond to the lateral boundary region, followed by the prognostic cells. The sorting of these prognostic cells with respect to their cell row stops after the first cell row (denoted by “sorted” vs. “unsorted” in Fig. 8.2).

Thus the indices are ordered in such a way that typical iterations over grid portions like prognostic cells, lateral boundary points etc. can be realized without conditional statements. Each portion is annotated by its start index, where the subscript corresponds to the \texttt{refin.cell ctrl} value. For convenience, there exists the auxiliary function \texttt{get.indices.c} (declared in the module \texttt{mo_loop_indices}) which helps to adjust the loop iteration accordingly: For a given value of \texttt{refin.c ctrl} and a specific block index we get the start and end indices to loop over.

Only the halo cells deserve some further remark (note the comment “\textit{when not overlapping with lateral boundary}“ in Fig. 8.2): In the special case, when no lateral boundary is present (global grid), the halo cells are stored in a contiguous fashion at the end of the index vector. When a lateral boundary is present, however, there exist some halo cells which also belong to the lateral boundary. These cells are then sorted into the leading part of the index array, since the ability to address boundary cells in a contiguous fashion is much more important in practice.
8.2. Data Structures

8.2.1. Description of the Model Domain: t_patch

The t_patch data structure contains all information about the grid coordinates and topology, as well as parallel communication patterns and decomposition information. It is declared in src/shr_horizontal/mo_model_domain.f90 as an array of length (# domains), where the coarsest base grid is denoted by the index 1, while the refined domains are denoted by numbers 2, 3 and so on.

All contained data arrays and indices relate to the index/block ordering described in Section 8.1 and non-existent indices are denoted by -1. The most important contents of the t_patch data structure are
The data members `cells`, `edges`, and `verts`, which are of the types `t_grid_cells`, `t_grid_edges`, and `t_grid_vertices`, respectively, give us information about the grid cells themselves, in particular about their geographical coordinates. For example,

```plaintext
t_grid_cells
  center(:,::) longitude and latitude of cell circumcenters, dimensions: [1:nproma, 1:nblks_c]
  neighbor_idx(:,::) line indices of triangles next to each cell, dimensions: [1:nproma, 1:nblks_c, 1:3]
  decomp_info information on domain decomposition :
```

Essentially, all data arrays which are contained in the grid files and which are described in Section 2.1.1 have a counterpart in this derived data type.

Besides, the data member `decomp_info` which separately exists for cells, edges and vertices, deserves additional comments. Its data type `t_grid_domain_decomp_info` is declared in `/src/parallel_infrastructure/mo_decomposition_tools.f90`:

```plaintext
t_grid_domain_decomp_info
  glb_index() global index of local cell, dimension: 1:n_patch_cells
  decomp_domain(:,::) domain decomposition flag, 0: owned (for cells), dimensions: [1:nproma, 1:nblks_c] :
```

The global index is particularly useful to perform operations (or write out data) which must not depend on the parallel domain decomposition of the model run. The “owner info” (`decomp_domain`) can be used to distinguish between prognostic cells and halo points whose values are just copied from adjacent PEs.
8.2.2. Date and Time Variables

When installing own processes within ICON’s time loop, the question for the current (simulation) time naturally arises. All global date and time variables are contained in the data structure `time_config`, which is of the derived data type `t_time_config` and declared in `src/configure_model/mo_time_config.f90`. The dates are initialized with the corresponding namelist parameters given in Section 5.1.1.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>tc_expstartdate</td>
<td>experiment start (tc means “time control”)</td>
</tr>
<tr>
<td>tc_expstopdate</td>
<td>experiment stop</td>
</tr>
<tr>
<td>tc_startdate</td>
<td>start of current simulation. In case of restart this is the date at which the simulation has been continued.</td>
</tr>
<tr>
<td>tc_stopdate</td>
<td>end of single run</td>
</tr>
<tr>
<td>tc_current_date</td>
<td>current model date</td>
</tr>
</tbody>
</table>

The dates and time spans make use of the `mtime` calendar library\(^2\) which is precise up to milliseconds without round-off errors. The `mtime` library resides in the directory `externals/mtime`. It is written in C and has a Fortran interface (module `mtime`).

We motivate the use of the `mtime` module by two examples. First, we perform a date calculation, adding a time span of 1 day to a given date. We make use of two variables: `mtime_date` (TYPE(datetime), POINTER) and `mtime_td` (TYPE(timedelta), POINTER).

```fortran
mtime_td => newTimedelta("P01D")
mtime_date => newDatetime("2014-06-01T00:00:00")

mtime_date = mtime_date + mtime_td
CALL datetimetostring(mtime_date, dstring)
WRITE (*,*) "2014-06-01T00:00:00 + 1 day = ", TRIM(dstring)
CALL deallocateDatetime(mtime_date)
CALL deallocateTimedelta(mtime_td)
```

As a second example, we demonstrate the `mtime` event mechanism which may be used to start certain processes in the program. An event (TYPE(event), POINTER) is defined by a start date, a regular trigger interval and an end date. Besides, let `RefDate` denote the event reference date (anchor date) in our example. Then the event triggers every `RefDate + k * interval`, but only within the bounds given by `startDate` and `endDate`.

```fortran
advectionEvent => newEvent('advection', RefDate, startDate, &
                          endDate, interval)
IF (isCurrentEventActive(advectionEvent, current_date)) THEN
```

---

\(^2\)The NWP mode uses the *proleptic* Gregorian calendar that is a backward extension of the Gregorian calendar to dates before its introduction October 15, 1582.
8.2 Data Structures

8.2.3. Data Structures for Physics and Dynamics Variables

On each model domain we need the same collection of 2D and 3D fields in order to describe the state of the atmosphere. These fields are collected in the data structure `t_nh_state`. This derived type and the following types are declared in `src/atm_dyn_iconam/mo_nonhydro_types.f90`.

First, the prognostic fields, which are integrated over time, are collected in the data structure `t_nh_prog`. Elements of `t_nh_prog` are allocated for each time slice that is needed for the time integration. For the nonhydrostatic time integration, the number of time slices is two, time `t` for the current time and `t + Δt` for the prediction.

<table>
<thead>
<tr>
<th><code>t_nh_prog</code></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>w</strong></td>
<td>orthogonal vertical wind [m s(^{-1})]</td>
</tr>
<tr>
<td><strong>vn</strong></td>
<td>orthogonal normal wind [m s(^{-1})]</td>
</tr>
<tr>
<td><strong>rho</strong></td>
<td>density [kg m(^{-3})]</td>
</tr>
<tr>
<td><strong>exner</strong></td>
<td>Exner pressure</td>
</tr>
<tr>
<td><strong>tke</strong></td>
<td>turbulent kinetic energy [m(^2) s(^{-2})]</td>
</tr>
<tr>
<td><strong>tracer</strong></td>
<td>tracer concentration [kg kg(^{-1})]</td>
</tr>
</tbody>
</table>

A global variable `p_nh_state` of type `t_nh_state` is instantiated in the module `/src/atm_dyn_iconam/mo_nonhydro_state.f90`. This is an array whose index corresponds to the model domain. The density of the atmosphere as state variable of the nonhydrostatic dynamical core is therefore given as

\[
p_{\text{nh state}}(\text{domain})\%\text{prog}(\text{time slice})\%\text{rho}(\text{index}, \text{level, block})
\]

Regarding the `time slice` argument, we briefly comment on ICON’s handling of the two-time-level scheme and the mechanism to avoid reallocation or unnecessary data copies:

For each prognostic variable in the two-time-level scheme, two arrays are pre-allocated:

\[
p_{\text{nh state}}\%\text{prog}(1)\%\text{field}(::,::)
\]
\[
p_{\text{nh state}}\%\text{prog}(2)\%\text{field}(::,::)
\]

Additionally, we introduce two global `INTEGER` index variables `nnow` and `nnew` which we initialize at model start with

\[
n\text{now} = 1
\]
\[
n\text{new} = 2
\]

The result values of the integration scheme (time slice `t + Δt`) are stored on the `nnew` time level. We therefore access the data on this time level by

\[
p_{\text{nh state}}\%\text{prog}(\text{nnew})\%\text{field}(::,::)
\]
While the calculations in the dynamical core fill this array with values, the prognostic state of the "old" time step can be accessed by

\[
p_{\text{nh \_state}}%\text{prog}(\text{nnow})\%\text{field}(\text{,:,:})
\]

Then, at the end of each dynamic (sub-)step, the time step \(n+1\) becomes the "old" one, while the time step \(n\) is freed and can be used as the new working array for the timestepping. This operation does not require any copying but merely exchanges the roles of \(\text{nnow}\) and \(\text{nnew}\):

\[
\text{CALL swap(nnow, nnew)}
\]

Alas, the whole process is complicated by the following two facts: First, \(\text{nnow}, \text{nnew}\) are defined for each domain separately. The above examples therefore require the domain index \(jg\), i.e. \(\text{nnow}(jg), \text{nnew}(jg)\). Second, as it has been explained in Section 3.7.1, different integration time steps are applied for efficiency reasons. A separate \(\text{nnow\_rcf}/\text{nnew\_rcf}\) accounting is required for the basic time step which is used for tracer transport, numerical diffusion and the fast-physics parameterizations, to distinguish it from the short time step used within the dynamical core\(^3\).

The data type \(\text{t\_nh\_diag}\) contains a collection of diagnostic fields, determined by all prognostic variables, boundary conditions and the compositions of the atmosphere.

<table>
<thead>
<tr>
<th>(\text{t_nh_diag})</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\text{u})  \  \ (\text{v})  \  \ (\text{temp})  \  \ (\text{pres})</td>
</tr>
<tr>
<td>zonal wind ([\text{m s}^{-1}])  \ meridional wind ([\text{m s}^{-1}])  \ temperature ([\text{K}])  \ pressure ([\text{Pa}])  \ :</td>
</tr>
</tbody>
</table>

Similar to the prognostic fields, the domain-wise data of type \(\text{t\_nh\_diag}\) can be accessed via \(p_{\text{nh \_state}(\text{domain})}\%\text{diag}\).

### 8.2.4. Parallel Communication

To simplify the data exchange between neighboring domain portions, ICON contains synchronization routines \text{exchange\_data}, defined in the module

```
src/parallel_infrastructure/mo_communication.f90
```

These take the specific halo region as an argument (data type \(\text{t\_comm\_pattern}\)) and several exchange patterns are pre-defined for each domain (see the derived data type \(\text{t\_patch}\)). For example, \(\text{comm\_pat\_c}\), defines the halo communication for \textit{cells} which are owned by neighboring processes. The subroutine call

\[
\text{CALL exchange\_data(patch\%comm\_pat\_c, array)}
\]

\(^3\)Here, the suffix \texttt{rcf} stands for “reduced calling frequency”.

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would perform a typical synchronization of the halo regions.

As it has been noted in Section 7.3.1, there exist different process groups in ICON: I/O, restarting and computation. These groups are related to MPI communicators which are defined in the module `src/parallel_infrastructure/mo_mpi.f90`. Probably the most important MPI communicator in the ICON code is `p_comm_work` (which is identical to the result of a call to `get_my_mpi_work_communicator()`). This is the MPI communicator for the work group.

The work group has a total size of `num_work_procs`, where each process may inquire about its rank by calling `get_my_mpi_work_id()`. On a non-I/O rank, its work group comprises all processes which take part in the prognostic calculations. It is therefore used by the `exchange_data` synchronization routine.

A final remark is related to the everlasting chit-chat of the status log (screen) output: The `process_mpi_stdio_id` is always the 0th process of the MPI communicator `process_mpi_all_comm`, which is the MPI communicator containing all PEs that are running this model component. A typical code line for printing out a message to screen would be

```fortran
IF (my_process_is_stdio()) write (0,*) "Hello world!"
```

With this, the message print-out would be suppressed on all PEs 1, 2, ...

Of course, there already exists an auxiliary subroutine `message()` in ICON, whose exact purpose is what we achieved manually above:

```fortran
CALL message('caller', 'message text')
```

The `message()` subroutine is located in the module `mo_exception` and restricts the print-out to PE #0. It takes the caller’s subroutine name as an additional argument.

### 8.3. Implementing Own Diagnostics

A thorough description of how to modify the ICON model and implement one's own diagnostics would certainly be a chapter in its own right. Here, we try to keep things as simple and short as possible with a view to the exercise Ex. C.8.3.

**Adding new fields.** ICON keeps so-called variable lists of its prognostic and diagnostic fields. This global registry eases the task of memory (de-)allocation and organizes the field’s meta-data, e.g., its dimensions, description and unit. The basic call for registering a new variable is the `add_var` command (module `mo_var_list`). Its list of arguments is rather lengthy and we will discuss them step by step.

First, we need an appropriate variable list to which we can append our new variable. For the sake of simplicity, we choose an existing diagnostic variable list, defined in the module `mo_nonhydro_state`:

```fortran
p_diag_list => p_nh_state_lists(domain)diag_list
```
The corresponding type definition can be found in the module `mo_nonhydro_types`. There, in the derived data type `TYPE(t_nh_diag)`, we place a 2D variable pointer

```fortran
REAL(wp), POINTER :: newfield(:, :)
```

which we can afterwards access as `p_nh_state(domain)%diag%newfield`.

Note that we did not allocate the variable so far.

Each ICON variable must be accompanied by appropriate meta-data. In this example we need to initialize GRIB and NetCDF variable descriptors for a variable located in the cell circumcenters (mass points). To keep this presentation as short as possible we have omitted the necessary USE statements:

```fortran
cf_desc = t_cf_var("newfield", "unit", "long name", DATATYPE_FLT32)
grib2_desc = grib2_var(discipline, parameterCategory, parameterNumber, &
                       DATATYPE_PACK16, GRID_UNSTRUCTURED, GRID_CELL)
```

The derived types `t_cf_var` and `t_grib2_var` are defined in the modules `mo_cf_convention` and `mo_grib2`, respectively. The expression `grib2_var` is actually a call to a constructor function, also defined in `mo_grib2`, which takes a triple of integers `(discipline, parameterCategory, parameterNumber)` as the field specifier.

Let us create an INTEGER array of length 2 with the name `shape2d_c`, denoting the dimensions of the new variable. The dimensions of a 2D field will be explained below. Here we take them as given:

```fortran
shape2d_c = (/ nproma, nblks_c /)
```

Now, with the essential ingredients at hand, we define our new field by the following call. We will place it at the very end of the subroutine `new_nh_state_diag_list` in the module `mo_nonhydro_state`.

```fortran
CALL add_var( p_diag_list, 'newfield',
              p_nh_state(domain)%diag%newfield,
              GRID_UNSTRUCTURED_CELL, ZA_SURFACE,
              cf_desc, grib2_desc,
              ldims=shape2d_c, lrestart=.FALSE. )
```

The INTEGER parameters `GRID_UNSTRUCTURED_CELL` and `ZA_SURFACE` define the type of the horizontal grid and the (trivial) vertical axis. From now on the new field can be specified in the output namelists that were described in Section 7.1:

```fortran
&output_nml
...
ml_varlist = 'newfield'
/
Looping over the grid points. Of course, the newly created field 'newfield' still needs to be filled with values. As explained in Section 8.1 above, nearly all DO loops that iterate over grid cells are organized in two nested loops: "jb loops" and "jc loops". Here the outer loop ("jb") is parallelized with OpenMP and limited by the cell block number nblks_c. The innermost loop iterates between 1 and the block length nproma.

Since the ICON model is usually executed in parallel, we have to keep in mind that each process can perform calculations only on a portion of the decomposed domain. Moreover, some of the cells between interfacing processes are duplicates of cells from neighboring sub-domains (so-called halo cells). Often it is not necessary to loop over these halo points, since they will be updated by the next parallel synchronization call.

The auxiliary function get_indices_c (declared in the module mo_loop_indices) helps to adjust the loop iteration accordingly:

\[
\begin{align*}
   i\_startblk &= p\_patch(domain)\%cells\%start\_block(grf\_bdywidth\_c+1) \\
   i\_endblk &= p\_patch(domain)\%cells\%end\_block(min\_rlcell\_int) \\
   &
\end{align*}
\]

\[
\begin{align*}
   DO \ j = i\_startblk, i\_endblk  \\
   \quad &CALL get\_indices\_c(p\_patch(domain), j, i\_startblk, i\_endblk, is, ie, gj, gie) \\
   &DO j = is, ie  \\
   \quad &p\_nh\_state(domain)\%diag\%newfield(jc,jb) = ...  \\
   END DO  \\
   END DO
\end{align*}
\]

The constants grf_bdywidth_c and min_rlcell_int can be found in the modules mo_Impl_constants_grf and mo_Impl_constants, respectively. Note that these constants have to inserted in start_block, end_block and also in the argument list of get_indices_c. A graphical interpretation of these constants is provided by Figure 8.2. The loop example therefore iterates over all prognostic cells (denoted by the blue area).

Loop exchange: The special preprocessor flag _LOOP_EXCHANGE can be found in numerous places of the ICON model code. It causes a loop interchange in many performance critical loops: If applied, the variable used in the inner loop switches to the outer loop.

Usually, this means that the loop over the vertical levels becomes the fastest running loop, compared to the iteration indices for the horizontal location. When arrays do not contain vertical levels, access to array elements may take advantage of the CPU cache.

Placing the subroutine call. Having encapsulated the computation together with the DO loops in a custom subroutine, we are ready to place this subroutine call in between ICON’s physics-dynamics cycle.

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Let us take a look at Figure 3.7: The outer loop “Dynamics → Physics → Output” is contained in the core module `mo_nh_stepping` inside the `TIME_LOOP` iteration. For diagnostic calculations it is important to have all necessary quantities available for input. On the other hand the result must be ready before the call to the output module,

```
CALL write_name_list_output(jstep)
```

The fail-safe solution here is to place the call immediately above this call.

Having inserted the call to the diagnostic field computation, we are done with the final step. Recompile the model code and you are finished!

**Style recommendations:** When writing your own extensions to ICON it is always a good idea to keep an eye on the quality of your code.

Make sure that there is no duplicate functionality and try to improve the readability of your subroutines through indentation, comments etc. This will make it easier for other developers to understand and assimilate. Better introduce own modules with complete interfaces and avoid USEs and PUBLIC fields.

A good starting point for your own project are the template files, given in the subdirectory `src/templates` of the ICON source code. These provide examples for modules, functions and subroutines.

**Additional remarks: 3D fields, accumulated quantities.** For the sake of brevity, only the simple case of two-dimensional fields has been discussed so far.

Three-dimensional fields would have an additional dimension for the column levels:

```
shape3d_c = (/ nproma, nlev, nblks_c /)
```

This information needs to be provided to the constructor `add_var(...)`, together with an INTEGER parameter which indicates the type of the vertical axis: For example, the pre-defined `ZA_HYBRID` would correspond to the most common category of 3D fields, the hybrid coordinates for the full levels (see Section 3.4).

Furthermore, it is often necessary to reset accumulated quantities in regular intervals. This can be achieved by

```
action_list = actions(new_action(ACTION_RESET, interval))
```

For example, by setting `interval = "PT06H"`, the respective field would be reset to zero every 6 hours.

```
CALL add_var( p_diag_list, 'newfield', &
 p_nh_state(domain)%diag%newfield, &
 GRID_UNSTRUCTURED_CELL, ZA_HYBRID, &
 cf_desc, grib2_desc, &
 action_list=actions(new_action(ACTION_RESET, interval)), &
 ldims=shape3d_c, lrestart=.FALSE. )
```
8.4. NWP Call Tree

All of ICON’s NWP and infrastructure modules, however numerous they may be, can roughly be classified into an initialization phase, the time integration loop and the clean-up phase. In Figure 8.3 we restrict ourselves to the most important subprograms.

These are listed together with a short description of their location and purpose, which, of course, change gradually between released versions. We recommend to compare this to the flow chart of processes in the physics-dynamics coupling, see Fig. 3.7.
8. Programming ICON

**Figure 8.3:** Call tree of ICON’s NWP component (note that this list has been restricted to the most important subroutines).
9. Post-Processing and Visualization

Oh my God! Look at that picture over there! There's the Earth coming up. Wow, is that pretty.

W. Anders, Apollo 8

ICON offers the possibility to produce output either in NetCDF or GRIB2 format. Many visualization environments such as GrADS, Matlab or R now include packages with which NetCDF data files can be handled. The GRIB format, which is also commonly used in meteorology, can be processed with these tools as well. However, since the standardization of unstructured GRIB records is relatively new, many post-processing packages offer only limited support for GRIB data that has been stored on the triangular ICON grid.

Since the visualization of regular (lat-lon) grid data is relatively straightforward, we will restrict ourselves in this course to a very simple program, ncview, which does not have a large functionality but is an easy-to-use program for a quick view of NetCDF output files. It is therefore very useful for a first impression.

Model data that has been stored on the triangular ICON grid can be visualized with the Python programming language, the NCL scripting language, R or the Generic Mapping Tools (GMT). Section 9.3 contains some examples how to visualize NetCDF data sets without the need of an additional regridding.

9.1. Retrieving Data Set Information

We begin with command-line utilities which provide a textual description of the data sets. For a quick overview of dimensions and variables, the command-line utility ncdump can be used. This program will shortly be described first. More sophisticated tools exist, for cutting out subsets of data, e.g., and producing averages or time series. One of these tools are the cdo utilities.

9.1.1. The ncdump Tool

The tool ncdump comes with the NetCDF library as provided by Unidata and generates a text representation of a NetCDF file on standard output. The text representation is in a form called CDL (network Common Data form Language). ncdump may be used as a simple browser for NetCDF data files, to display the dimension names and sizes, variable names, types and shapes, attribute names and values, and optionally, the data values.
themselves for all or selected variables in ASCII format. For example, to investigate the structure of a NetCDF file, use

```
ncdump -h data-file.nc
```

Dimension names and sizes, variable names, dependencies and values of dimensions will be displayed. To show the type version of a NetCDF file, type

```
ncdump -k data-file.nc
```

This gives information about the NetCDF base format of a specific file, for example NetCDF Classic Format, NetCDF 64-bit Offset Format or NetCDF-4 Format.

NetCDF data can also be redirected to a text file with

```
ncdump -b c data-file.nc > data-file.cdl
```

This produces an annotated CDL version of the structure and the data in the NetCDF file `data-file.nc`. You can also save data for specified variables to text files just using:

```
ncdump -v variable data-file.nc > data-file.txt
```

For further information on working with `ncdump` see

http://www.unidata.ucar.edu/software/netcdf/docs/netcdf_utilities_guide.html#ncdump_guide

### 9.1.2. CDO – Climate Data Operators

The CDO (Climate Data Operators) are a collection of command-line operators to manipulate and analyze NetCDF and GRIB data. The CDO package is developed and maintained at MPI for Meteorology in Hamburg. Source code and documentation are available from

https://code.mpimet.mpg.de/projects/cdo

The tool includes more than 400 operators to print information about data sets, copy, split and merge data sets, select parts of a data set, compare data sets, modify data sets, arithmetically process data sets, to produce different kind of statistics, to detrend time series, for interpolation and spectral transformations. The CDOs can also be used to convert from GRIB to NetCDF or vice versa, although some care has to be taken there.

In particular, the "operator" `cdo infov` writes information about the structure and contents of all input files to standard output. By typing

```
cdo infov data-file.nc
```

in the command-line for each field the following elements are printed: date and time, parameter identifier and level, size of the grid and number of missing values, minimum, mean and maximum. A variant of this CDO operator is

```
cdo sinfov data-file.nc
```

which prints out short information of each field.
Remapping to Regular Grids with CDO. The CDO are, by the way, also capable of remapping data to regular grids. Basically, two steps are necessary to process ICON output:

- In the first call of the CDO ("gencon"), coefficients are generated and stored to a separate file:

  \[ \texttt{cdo gencon, lat-lon\_grid\_description icon\_grid coefficient\_file} \]

  The generation of the interpolation weights may take some time.

- Afterwards, the interpolation is done with

  \[ \texttt{cdo remap, grid\_description, coefficient\_file in\_file out\_file} \]

Here, the file \texttt{icon\_grid} is a NetCDF file which contains the topological and geometric information about the triangular ICON grid. This grid information must correspond to the data set \texttt{in\_file} and it is not part of the data set itself. Instead, it must be downloaded separately from the web. See \url{http://icon-downloads.mpimet.mpg.de/dwd_grids.xml} for the list of "official" DWD ICON grids.

The structure of the lon-lat grid description file is explained in the CDO documentation\(^\text{2}\). A global regular grid, for example, would be defined as

\begin{verbatim}
gridtype = lonlat
xsize   = 1440
ysize   = 721
xfirst  = 0.00
xinc    = 0.25
yfirst  = -90.00
yinc    = 0.25
\end{verbatim}

Caveat: Alas, the CDO do not handle the entire set of GRIB2 meta-data correctly. Some meta-data items, for example those which regard ensemble runs, still remain unsupported. However, this limitation does not matter if the desired output format is NetCDF and not GRIB2, or if the processed data fields are rather standard. Alternatives for remapping ICON data sets to lon-lat grids are the "fieldextra" software (which is a COSMO software) or the DWD ICON Tools, which are internally used at DWD but lack official support.

9.2. Plotting Data Sets on Regular Grids: \texttt{ncview}

\texttt{ncview} is a visual browser for NetCDF format files developed by David W. Pierce. Using \texttt{ncview} you can get a quick and easy look at regular grid data in your NetCDF files.

\textit{Note for the DWD training course that the visualization tools (CDO, NCL, \texttt{ncview}) which are described in this section are only available on the \url{ice.dwd.de}!}

\(^{\text{1}}\)Currently in operational use is the global grid #26 with 13 km horizontal grid spacing (\url{http://icon-downloads.mpimet.mpg.de/grids/public/edzw/icon_grid_0026_R03B07_G.nc}).

\(^{\text{2}}\)Appendix D, \url{https://code.mpimet.mpg.de/projects/cdo/embedded/index.html}.\n
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To install ncview on your local platform, see the ncview website:
http://meteora.ucsd.edu/~pierce/ncview_home_page.html

You can run the program by typing:

ncview data-file.nc

which will open a new window with the display options. It is possible to view simple movies of data, view along different dimensions, to have a look at actual data values at specific coordinates, change color maps, invert data, etc., see the screenshot in Fig. 9.1.

If data-file.nc contains wildcards such as ‘*’ or ‘?’ then all files matching the description are scanned, if all of the files contain the same variables on the same grid. Choose the variable you want to view. Variables which are functions of longitude and latitude will be displayed in two-dimensional images. If there is more than one time step available you can easily view a simple movie by just pushing the forward button. The appearance of the image can be changed by varying the colors of the displayed range of the data set values or by adding/removing coastlines. Each one- or two-dimensional subset of the data can be selected for visualization. ncview allows the selection of the dimensions of the fields available, e.g. longitude and height instead of longitude and latitude of 3D fields.

The pictures can be sent to Postscript (*.ps) output by using the function print. Be careful that whenever you want to close only a single plot window to use the close button, because clicking on the ⊠-icon on the top right of the window will close all ncview windows and terminate the entire program!

9.3. Plotting Data Sets on the Triangular Grid

Let us now focus on ICON’s original computational mesh, the triangular grid. In this section we will sketch several approaches for the visualization of data sets based on the triangular cells.
By now there is quite a number of scripting languages and visualization tools available for this task. We will present quick start examples for the Python programming language, NCL, R, and GMT. All these approaches have in common that the NetCDF grid file must be read in together with the data file.

9.3.1. Visualization with Python

The Python module PyNGL (pronounced “pingle”) is a Python language module for generating high-quality, 2D visualizations of scientific data. PyNIO is a Python module used for reading and writing NetCDF, GRIB, and HDF.

PyNGL is developed by the Computational and Information Systems Lab at the National Center for Atmospheric Research (NCAR), see

https://www.pyngl.ucar.edu

In the following we will explain the basic usage of PyNGL in a step-by-step tutorial.

PyNGL Installation

PyNGL and PyNIO can be installed on Linux and MacOS systems via conda\(^3\), or its slimmed-down version miniconda:

First, download the miniconda installer script to your machine. In our example we will install the stable release version for Linux x86 architectures. Visit the conda web page to find out which version is the right one for your platform.

```
wget https://repo.anaconda.com/miniconda/Miniconda3-latest-Linux-x86_64.sh
bash Miniconda3-latest-Linux-x86_64.sh
source /e/uhome/fprill/miniconda3/etc/profile.d/conda.sh
```

Once miniconda is installed, we use the conda command to install the PyNGL and PyNIO Python packages:

```
conda create --name pyn_env --channel conda-forge pyngl
conda install --name pyn_env --channel conda-forge pynio
```

Now, you can activate Python environment via

```
conda activate pyn_env
```

and the newly installed packages PyNGL and PyNIO are ready to use.

\(^3\) Conda is an open source package management system see https://conda.io.
PyNGL Step-by-step Tutorial

In the following we provide a detailed step-by-step tutorial for producing graphics from an ICON data set. To this end, we create a file `visualization_tutorial.py` where a sequence of commands can be stored and executed with

```python
python visualization_tutorial.py
```

We begin by loading some Python modules which provide high-level functions for plotting and numerical computations.

```python
# load Python modules for plotting
import Ngl, Nio, numpy
```

These lines also contain a comment. Comment lines in Python are preceded by the hash sign `#`.

**Step 1: Reading Grid Coordinates From File.** Since the ICON model uses an unstructured grid topology, we open and read such a topology file, stored in NetCDF format, by the following commands:

```python
gridfile = Nio.open_file("gridfile.nc")
print(gridfile)
```

The `print` command lists all variables that have been found in the NetCDF file as textual output. For the ICON grid, the vertex positions of the grid triangles are of special interest. They are stored as longitude/latitude positions in the `vlon`, `vlat` fields (this is explained in more detail in Section 2.1.1, page 19). For PyNGL we convert from steradians to degrees:

```python
vlon = numpy.rad2deg( gridfile.variables["vlon"][:]
vlat = numpy.rad2deg( gridfile.variables["vlat"][:]
```

Additionally, we load the vertex indices for each triangle edge of the icosahedral mesh.

```python
edge_vertices = gridfile.variables["edge_vertices"][:]
```

The indices are stored in the grid file data set `edge_vertices` and reference the corresponding vertices from `vlon`, `vlat`,

```python
data #i: (vlon[q1],vlat[q1]) — (vlon[q2],vlat[q2])
```

where

```python
q1/2 := edge_vertices[1/2,i] − 1
```

Note that by subtracting 1 we take the 0-based array indexing of Python into account.
Step 2: Creating a Plot of the Triangular Grid. Producing graphics with PyNGL requires the creation of a so-called workstation, i.e. a description of the output device. In this example, this “device” will be an image file plot.png, but we could also define a different output format, e.g. PostScript "ps" instead.

```
   wks = Ngl.open_wks("png","plot")
```

Then the map settings have to be defined and we collect these specifications in a “resource data structure” named `config1`. First of all, we disable the immediate drawing of the map image, since the ICON icosahedral grid plot will consist of two parts: the underlying map and the grid lines. We do so by setting the resource attributes `bglFrame` and `nglDraw` to `False`.

We then define an orthographic projection centered over Europe. It is important that grid lines are true geodesic lines, otherwise the illustration of the ICON grid would contain graphical artifacts, therefore we set the parameter `mpGreatCircleLinesOn`.

```
   config1 = Ngl.Resources()
   config1.nglFrame = False
   config1.nglDraw = False
   config1.mpProjection = "Orthographic"
   config1.mpGreatCircleLinesOn = True
   config1.mpCenterLonF = 10
   config1.mpCenterLatF = 50
```

Having completed the setup of the `config1` data structure, we can create an empty map by the following command:

```
   map = Ngl.map(wks,config1)
```

Now, the edges of the ICON grid must be added to the plot. As described before, we convert the indirectly addressed `edge_vertices` into an explicit list of geometric segments with dimension $2 \times nedges$:

```
   ecx = numpy.ravel( vlon[ edge_vertices-1 ], order='F' )
   ecy = numpy.ravel( vlat[ edge_vertices-1 ], order='F' )
```

This operation deserves additional comments: First of all, since the PyNGL plotting function below will expect one-dimensional lists for its interface, we use the auxiliary function `numpy.ravel` for reshaping the array of lines. Second, with `order='F'` we define the array ordering of `vlon`, `vlat` to be “Fortran-style”. Compared to Python (numpy) this is the transpose memory layout.

Now, there exists an PyNGL high-level command for plotting lines, `Ngl.add_polyline`. 
Figure 9.2.: The two plots generated by the Python example script in Section 9.3.1.

```python
lines_cfg = Ngl.Resources()
lines_cfg.gsSegments = numpy.arange(0, edge_vertices.size, 2)
poly = Ngl.add_polyline(wks, map, ecx, ecy, lines_cfg)
```

The whole plotting process is then triggered by the command

```python
Ngl.draw(map)
Ngl.frame(wks)
Ngl.end()
```

Here, the call to `Ngl.end()` terminates the PyNGL script. The first of the resulting image files `plot.000001.png` will contain an illustration similar to Fig. 9.2 (left part).

**Step 3: Loading a Data Set from a Second File.** We now assume that the data sets produced by the ICON model have been stored in NetCDF format. This allows to visualize the unstructured data without additional Python packages. As a second file we open such a NetCDF data set `datafile.nc` in read-only mode and investigate its data set `topography_c`:

```python
datafile = Nio.open_file("datafile.nc")
topo = datafile.variables["topography_c"]
print(topo)
```

The final step of this exercise is the creation of a contour plot from the data contained in `datafile`. As it has been stated by the previous call to `print`, the data sites for the field `topography_c` are the triangle circumcenters, located at `clon`, `clat`.

```python
clon = numpy.rad2deg( gridfile.variables["clon"][::] )
clat = numpy.rad2deg( gridfile.variables["clat"][::] )
```
For a basic contour plot, a cylindrical equidistant projection with automatic adjustment of contour levels will do. It is important to specify the two additional arguments `sfXArray` and `sfYArray`.

```python
config2 = Ngl.Resources()
config2.mpProjection = "CylindricalEquidistant"
config2.cnFillOn = True
config2.cnLinesOn = False
config2.cnLineLabelsOn = False
config2.sfXArray = clon
config2.sfYArray = clat
```

Afterwards, we generate the plot (image file `plot.000002.png`) with a call to `Ngl.contour_map`.

```python
map = Ngl.contour_map(wks,topo,config2)
```

Note that this time it is not necessary to launch additional calls to `draw` and `frame`, since the default options in `config2` are set to immediate drawing mode.

You may wonder why the plot has a rather smooth appearance without any indication of the icosahedral triangular mesh. What happened is that PyNGL generated its own Delaunay triangulation building upon the cell center coordinates provided via `clon`, `clat`. Thus, we are unable to locate and investigate individual ICON grid cells. In order to visualize individual cells, we need to additionally load the vertex coordinates of each triangle into PyNGL. This information is also available from the grid file and is stored in the fields `clon_vertices`, `clat_vertices`.

```python
clonv = numpy.rad2deg( gridfile.variables["clon_vertices"][:,:] )
clatv = numpy.rad2deg( gridfile.variables["clat_vertices"][:,:] )
clon_vertices = clonv
clat_vertices = clatv
config2.sfXCellBounds = clon_vertices
config2.sfYCellBounds = clat_vertices
config2.cnFillMode = "CellFill"
```

By choosing the `CellFill` mode, it is ensured that every grid cell is filled with a single color.

Afterwards we generate the plot once more with a call to `Ngl.contour_map`.

```python
map = Ngl.contour_map(wks,topo,config2)
```

Do you see the difference?
9.3.2. NCL – NCAR Command Language

The NCAR Command Language (NCL) is an interpreted language designed specifically for scientific data analysis and visualization. It is built on top of the same “resource model” used in the PyNGL Python package.

NCL allows convenient access to data in a variety of formats such as NetCDF and GRIB1/2, among others. It has many features common to modern programming languages, such as types, variables, operators, expressions, conditional statements, loops, and functions and procedures, see https://www.ncl.ucar.edu for details.

Besides an interactive mode, NCL allows for script processing (recommended). NCL scripts are processed on the command-line by typing

```
ncl filename.ncl
```

For visualizing ICON data on the native triangular grid, we recommend using NCL 6.2.0 or higher.

Note that NCAR has recently made the decision to adopt Python as the scripting language platform of choice for future development of analysis and visualization tools. NCL’s graphics routines will have continued development through the PyNGL Python package.

**NCL Quick-Start Example**

The following example script creates a temperature contour plot with NCL (see Figure 9.3):

```
begin
  ; Open model level output file
  File = addfile( "JABW_DOM01_ML_0001.nc", "r" )

  ; read grid information (i.e. coordinates of cell centers and vertices)
  rad2deg = 45./atan(1.) ; radians to degrees
  clon = File->clon * rad2deg ; cell center, lon (ncells)
  clat = File->clat * rad2deg ; cell center, lat (ncells)
  vlon = File->clon_bnds * rad2deg ; cell vertices, lon (ncells,3)
  vlat = File->clat_bnds * rad2deg ; cell vertices, lat (ncells,3)

  ; read data
  temp_ml = File->temp(:,,:,:); dims: (time,lev,cell)
  print("max T " + max(temp_ml) )
  print("min T " + min(temp_ml) )

  ; create plot
  wks = gsn_open_wks("ps","outfile")
  gsn_define_colormap(wks,"testcmap") ; choose colormap
```

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To open a data file for reading, the function `addfile` returns a file variable reference to the specified file. Second, for drawing graphics, the function `gsn_open_wks` creates an output resource, where the “ps”, “pdf” or “png” format are available. Third, the command `gsn_csm_contour_map` creates and draws a contour plot over a map.

Loading the coordinates of the triangle cell centers into NCL (resources `sfXArray` and `sfYArray`) is essential for visualizing ICON data on the native grid. Loading the vertex coordinates of each triangle (resources `sfXCellBounds` and `sfYCellBounds`), however, is optional. If not given, a Delaunay triangulation will be performed by NCL, based on the cell center information. If given, the triangles defining the mesh will be deduced by sorting and matching vertices from adjacent cell boundaries. If you are interested in the correct representation of individual cells, the resource `sf[X/Y]CellBounds` should be set.

Creating a plot can get very complex depending on how you want to look at your data. Therefore we refer to the NCL documentation that is available online under

![Figure 9.3: ICON temperature field on a specific model level produced with the above NCL script.](image)
Note that there is another NCL example contained in the course material, which very closely resembles the steps explained in the PyNGL tutorial in Section 9.3.1. For the exercises in this tutorial we refer to the prepared NCL scripts. These files are stored in the subdirectory test_cases/casexx together with the model run scripts.

9.3.3. Visualization with R

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R is a free software environment for statistical computing and graphics. R is available as free software under the terms of the Free Software Foundations GNU General Public License. More Information can be found here:

https://www.r-project.org

To start R in an interactive mode simply type

R

on the command line. Afterwards R commands can be entered, which are then interpreted line by line. R can be extended (easily) via packages. Additional packages have to be installed via the R command

install.packages("package_name")

For the compilation of some packages it might be necessary to provide specific (development) libraries on the system and to give information about the include and library paths as additional arguments to that command.

Most packages are available through the CRAN family of internet sites. An exception to this is the gribr package, which is used in the example script below to read in GRIB2 data:

https://github.com/nawendt/gribr

This package uses and therefore needs a recent installation of ecCodes (it is not working with the GRIB API). Additional information about (other) prerequisites and the installation of the package can be found on the given website.

Besides an interactive mode, R allows for script processing (recommended). R scripts are processed on the command-line by typing

Rscript filename.R

As default a PDF named Rplots.pdf will be created.
Figure 9.4.: ICON 2m temperature field produced with the given example script for R.

R Quick-Start Example

The following example script `icon_native4_tutorial.R` can be found in the tarball directory `scripts`. It creates a global temperature contour plot with R (see Figure 9.4):

```
# ... on LCE, as a prerequisite issue the following module commands
# module unload oracle R eccodes
# module load unsupported oracle/18.3.0.0 R/3.5.1
# module load eccodes/2.9.0 eccodes/R-gribr-2.9.0-prerelease
#
# load necessary libraries
library(gribr)
library(RNetCDF)
library(data.table)
library(ggplot2)
library(dplyr)
library(colorRamps)
#
# --- setup -----------------------------------------------

shortName <- "T_2M"
title <- paste(shortName, " ICON (R2B6)\n")
```
# data file names
grid <- "/icon_grid_0024_R02B06_G.nc"
data <- "/T_2M.R2B06_ICON_global.grb"

# --- icon grid ------------------------------------------------
ncHandle <- open.nc(grid)

# function to convert coordinates in degrees
rad2deg <- function(rad) {(rad * 180) / (pi)}

# get longitudes and latitudes of triangle vertices of a cell
vlon <- rad2deg(var.get.nc(ncHandle,"clon_vertices"))
vlat <- rad2deg(var.get.nc(ncHandle,"clat_vertices"))

close.nc(ncHandle)

# --- icon data ------------------------------------------------
gribHandle <- grib_open(data)

# select the grib record based on a given list of keys
gribRecord <- grib_select(gribHandle, list(shortName = shortName))

grib_close(gribHandle)

# create a data table with data to plot - ids and values are tripled
DT <- data.table(lon = as.vector(vlon),
                 lat = as.vector(vlat),
                 id = rep(1:(dim(vlon)[2]) , each=3),
                 var = rep(gribRecord$values, each=3))

# --- domain and edges -----------------------------------------

# Global
xrange <- c(-180., 180.)
yrange <- c(-90., 90.)

# select the subset of ids in the domain
usedIDs <- unique(DT[lon%between%xrange & lat%between%yrange]$id)

# use only the respective subset of the data
DT <- DT[id %in% usedIDs]

# special treatment for the cells near the date line
# . . . when the 3 corners on opposite sides of the date line move one corner
IDsR = DT[,list( (max(lon)-min(lon))>200 & mean(lon)>0.0 ), by=id][V1==T]$id
IDsL = DT[,list( (max(lon)-min(lon))>200 & mean(lon)<0.0 ), by=id][V1==T]$id
DT[id %in% IDsR & lon < 0.0, lon := lon + 360.0]
DT[id %in% IDsL & lon > 0.0, lon := lon - 360.0]

# . . . copy triangles by 360deg to fill holes near date line
DTT <- DT[id%in%IDsR]
DTT[lon > 0.0, lon := lon - 360.0]
9.3 Plotting Data Sets on the Triangular Grid

```r
# Visualization

9.3.4. GMT – Generic Mapping Tools

GMT is an open source collection of command-line tools for manipulating geographic and Cartesian data sets and producing PostScript illustrations ranging from simple x-y plots via contour maps to 3D perspective views. GMT supports various map projections and transformations and facilitates the inclusion of coastlines, rivers, and political boundaries. GMT is developed and maintained at the University of Hawaii, and it is supported by the National Science Foundation.

To install GMT on your local platform, see the GMT website:

http://gmt.soest.hawaii.edu

Since GMT is comparatively fast, it is especially suited for visualizing high resolution ICON data on the native (triangular) grid. It is capable of visualizing individual grid cells
and may thus serve as a helpful debugging tool. So far, GMT is not capable of reading ICON NetCDF or GRIB2-output offhand. However, CDO can be used to convert your data to a format readable by GMT.

From your NetCDF output, you should first select your field of interest and pick a single level at a particular point in time:

```
cdo -f nc selname,VNAME -seltimestep,ITIME -sellevidx,ILEV 
   ICON_OUTPUT.nc ICON_SELECTED.nc
```

Now this file must be processed further using the `outputbounds` command from CDO, which finally leads to an ASCII file readable by GMT.

```
cdo -outputbounds ICON_SELECTED.nc > ICON_SELECTED.gmt
```

The output looks as follows:

```plaintext
# Generated by CDO version 1.6.3
#
# Operator = outputbounds
# Mode = horizontal
#
# File = NWP_DOM01_ML_0006_temp.nc
# Date = 2012-01-04
# Time = 00:00:00
# Name = temp
# Code = 0
# Level = 80
#
> -Z254.71
   -155.179  90
   36   89.5695
   108   89.5695
   -155.179  90
> -Z255.276
   36   89.5695
   36   89.1351
   72   89.2658
   36   89.5695
...
```

For each triangle, it contains the corresponding data value (indicated by `-Z`) and vertex coordinates.

As a starting point, a very basic GMT script is added below. It visualizes the content of `test.gmt` on a cylindrical equidistant projection including coastlines and a colorbar. An example plot based on this script is given in Figure 9.5.

```bash
#!/bin/bash

# Input filename
```
9.3 Plotting Data Sets on the Triangular Grid

```plaintext
INAME="test.gmt"

# Output filename
ONAME="test.ps"

# generate color palette table (min/max/int)
makecpt -Cpolar -T"235"/"305"/"5" > colors.cpt

# draw triangle and take fill color from colors.cpt
psxy ${INAME} \\
-Rd -Jq0/1:190000000 -Ccolors.cpt -X3.2 -Y4. -K > ${ONAME}

# visualize coastlines
pscoast -Rd -Jq0/1:190000000 -Dc -W0.25p,black -K -O >> $ONAME

# plot colorbar
psscale -D11c/14c/18c/1.0ch -Ccolors.cpt -E -B:"T":/:K: -U -O >> $ONAME
```

Note: In order to get filled polygons, the -L option must be added to psxy. The purpose of -L is to force closed polygons, which is a prerequisite for polygon filling. However, in the latest release of GMT (5.1.2) adding this option results in very large output files whose rendering is extremely slow. Thus, the -L option was omitted here so that only triangle edges are drawn and colored.

Figure 9.5.: ICON temperature field on a specific model level produced with the above GMT script.
10. Running ICON-ART

In this lesson you will learn how to run the package for Aerosols and Reactive Trace Gases, ICON-ART (Rieger et al., 2015, Schröter et al., 2018).

10.1. General Remarks

ICON-ART is an extension of the ICON model that was developed at the Institute of Meteorology and Climate Research (IMK) at the Karlsruhe Institute of Technology (KIT). It allows the online calculation of reactive trace substances and their interaction with the atmosphere. The interfaces to the ART code are part of the official ICON code.

In order to obtain the ART code, the institution that wants to use ICON-ART has to sign an additional license agreement with Karlsruhe Institute of Technology (KIT). Further information can be found on the following website:

http://icon-art.imk-tro.kit.edu

After you have signed the license agreement, you will be provided with a compressed file with the recent source code of ART which is called ART-v<X>,<YY>.tar.gz. <X> and <YY> indicate the version number.

10.2. ART Directory Structure

The ART directory contains several subdirectories. The purposes of those subdirectories are explained in the following.

The Directory aerosol_dynamics

ICON-ART solves the diffusion equation of aerosol. For this purpose, the following processes have to be considered: Advection$^1$, turbulent diffusion$^1$, changes due to subgrid-scale convective transport$^1$, sedimentation, washout, coagulation, condensation from the gas-phase, radioactive decay, and emissions$^2$. With a few exceptions (marked by $^1$ and $^2$), the modules calculating the tendencies due to these processes are stored within the aerosol_dynamics folder. Additionally, routines calculating diagnostic properties that are needed as input for the aerosol process parameterizations are stored within this folder. The exceptions are the following:
• 1: The tendencies due to these processes are calculated within the ICON code. This is part of the tracer framework of ICON.

• 2: The emission routines are an important source for atmospheric aerosol. ART offers the option to easily plug in new emission schemes. In order to keep clarity within the folders, emissions routines get their own folder emissions (see below).

The Directory chemistry

ICON-ART solves the diffusion equation of gaseous tracers. Besides advection, turbulent diffusion and subgrid-scale convective transport which are treated by the ICON tracer-framework, this includes also chemical reactions. The chemistry directory contains the routines to calculate chemical reaction rates of gaseous species.

The Directory emissions

Within the emissions directory, emission routines for aerosol and gaseous species are stored.

The Directory externals

Within the externals directory, code from external libraries is stored (i.e. cloud, mecicon).

The Directory io

The io directory contains input and output routines.

The Directory chem.init

The chem.init directory contains routines needed for an initialization of ICON-ART tracers with Mozart results.

The Directory phy.interact

Modules within the phy.interact directory treat the direct interaction of aerosol particles and trace gases with physical parameterizations of ICON. Examples are the interaction of aerosols with clouds (i.e. the two-moment microphysics) and radiation.
The Directory runctrl_examples

The runctrl_examples directory contains the following subdirectories

**emiss_ctrl** storage of example emission files for volcanic emissions, radioactive release

**init_ctrl** location of coordinate file for MOZART initialisation and initialisation table for LINOZ (linearized ozone) algorithm

**photo_ctrl** location of CloudJ Input files (Cross sections, Q-Yields)

**run_scripts** runscripts for testsuite and training course testcases

**xml_ctrl** storage of basic .xml files for tracer registration and system files (.dtd)

The Directory shared

The shared directory contains a collection of routines that do not fit into other categories. This applies mostly to initialization and infrastructure routines.

The Directory tools

The tools directory contains helpful tools for ART developers (e.g. a generalized clipping routine).

10.3 Installation

In this section, a brief description of how to compile ICON-ART is given. The user has to do the same steps as compiling ICON with a few additions. The reader is referred to Section 1.2.2 in order to compile ICON successfully. First, the `art.tar.gz` file has to be uncompressed. You will obtain a directory, which should be copied inside the ICON source directory `$ICON-DIR/src/`. In the following, we refer to this directory `$ICON-DIR/src/art` as `$ARTDIR`.

If you have compiled ICON without ART before, you have to do clean up first:

```
make distclean
```

In order to compile ICON-ART, an additional flag has to be set at the configuration command:

```
./configure --with-fortran=cray --with-art
```

By setting `--with-art` a compiler flag `-D_ICON_ART` is set. This flag tells the preprocessor to compile the code inside the ART interfaces and hence connect the ICON code with the ART code. As soon as the configuration is finished, you can start to compile the ICON-ART code:

```
./build_command
```
10.4. Configuration of an ART Job

10.4.1. Recommended ICON Namelist Settings

It is necessary for the user to choose the ICON settings carefully to obtain a stable ICON-ART simulation with scientifically reasonable results. Hence, the user should pay special attention to the namelist parameters listed in table 10.1.

Table 10.1.: Recommended ICON namelist settings for ART tracers.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Namelist</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dtime</td>
<td>-</td>
<td>run_nml</td>
<td>If facing stability problems, it is recommended to use a shorter time step as recommended by operational setups (e.g. ( \frac{3}{4} ) dtime_{oper}).</td>
</tr>
<tr>
<td>ndyn_substeps</td>
<td>-</td>
<td>run_nml</td>
<td>There is no need to call the dynamics more often than in operational setup. Adjust it according to your dtime choice.</td>
</tr>
</tbody>
</table>

10.4.2. ART Namelists

ICON-ART has an own namelist to modify the setup of ART simulations at runtime. The main switch for ART, lart, is located inside run_nml. The namelist for the other ART switches is called art_nml.

A naming convention is used in order to represent the type of data. An INTEGER namelist parameter starts with iart, a REAL namelist parameter start with rart, a LOGICAL namelist parameter starts with lart, and a CHARACTER namelist parameter starts with cart.

The ICON-ART namelist is located in the module src/namelists/mo_art_nml.f90. General namelist parameters are listed and explained within Table 10.2. Namelist parameters for ART input are listed within Table 10.3. Namelist parameters related to atmospheric chemistry are listed within Table 10.4. Namelist parameters related to aerosol physics are listed within Table 10.5.
### Table 10.2.

General namelist parameters to control the ART routines. These switches are located inside `art.nml`. The only exception is the `lart` switch which is located in the `run.nml`.

<table>
<thead>
<tr>
<th>Namelist Parameter</th>
<th>Default</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>lart</code></td>
<td>.FALSE.</td>
<td>Main switch which enables the ART modules. Located in the namelist <code>run.nml</code>.</td>
</tr>
<tr>
<td><code>lart_chem</code></td>
<td>.FALSE.</td>
<td>Enables chemistry. The chemical mechanism and the according species are set via <code>iart_chem_mechanism</code>.</td>
</tr>
<tr>
<td><code>lart_pntSrc</code></td>
<td>.FALSE.</td>
<td>Enables point sources for passive tracer. The sources are controlled via <code>cart_pntSrc_xml</code>. See also Section 10.4.5.</td>
</tr>
<tr>
<td><code>lart_aerosol</code></td>
<td>.FALSE.</td>
<td>Main switch for the treatment of atmospheric aerosol.</td>
</tr>
<tr>
<td><code>lart_passive</code></td>
<td>.FALSE.</td>
<td>Main switch for the treatment of passive tracer.</td>
</tr>
<tr>
<td><code>lart_diag_out</code></td>
<td>.FALSE.</td>
<td>If this switch is set to .TRUE., diagnostic output fields are available. Set it to .FALSE. when facing memory problems.</td>
</tr>
</tbody>
</table>

### Table 10.3.

Namelist parameters to control ART input. These switches are located inside `art.nml`. For details regarding the tracer and modes initialization with XML files, see Section 10.4.3.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>cart_chemistry_xml</code></td>
<td>''</td>
<td>Path and file name to the XML file for specifying chemical tracer. See also Section 10.4.3.</td>
</tr>
<tr>
<td><code>cart_aerosol_xml</code></td>
<td>''</td>
<td>Path and file name to the XML file for specifying aerosol tracer. See also Section 10.4.3.</td>
</tr>
<tr>
<td><code>cart_passive_xml</code></td>
<td>''</td>
<td>Path and file name to the XML file for specifying passive tracer. See also Section 10.4.3.</td>
</tr>
<tr>
<td><code>cart_modes_xml</code></td>
<td>''</td>
<td>Path and file name to the XML file for specifying aerosol modes. See also Section 10.4.4.</td>
</tr>
<tr>
<td><code>cart_pntSrc_xml</code></td>
<td>''</td>
<td>Path and file name to the XML file for specifying point source emissions. See also Section 10.4.5.</td>
</tr>
</tbody>
</table>
Table 10.4.: Namelist parameters related to atmospheric chemistry. These switches are located inside `art_nml`.

<table>
<thead>
<tr>
<th>Namelist Parameter</th>
<th>Default</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>iart_chem_mechanism</td>
<td>0</td>
<td>Sets the chemical mechanism and takes care of the allocation of the according species. Possible values:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0: Stratosph. short-lived Bromocarbons, 1: Simplified OH chemistry, takes photolysis rates into account</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2: Full gas phase chemistry</td>
</tr>
</tbody>
</table>

Table 10.5.: Namelist parameters related to aerosol physics. These switches are located in `art_nml`.

<table>
<thead>
<tr>
<th>Namelist Parameter</th>
<th>Default</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>iart_seasalt</td>
<td>0</td>
<td>Treatment of sea salt aerosol. Possible values:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0: No treatment.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1: As specified in Lundgren et al. (2013).</td>
</tr>
<tr>
<td>iart_volcano</td>
<td>0</td>
<td>Treatment of volcanic ash particles. Possible values:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0: No treatment.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2: 2-moment treatment.</td>
</tr>
<tr>
<td>cart_volcano_file</td>
<td>''</td>
<td>Path and filename of the input file for the geographical positions and the types of volcanoes.</td>
</tr>
</tbody>
</table>
10.4.3. Tracer Definition with XML Files

The definition of tracers in ICON-ART is done with the use of three XML files. A distinction between aerosol, chemical and passive tracers is made. Aerosol tracers are defined by an XML file specified with the namelist switch cart_aerosol_xml containing all liquid and solid particles participating in aerosol dynamics. Chemical tracers are defined by an XML file via cart_chemistry_xml and cover gaseous substances participating in chemical reactions. Passive comprises all gaseous, liquid and solid tracers that are only participating in transport processes. These are specified with the XML file cart_passive_xml. With the following example XML file, two passive tracers called trPASS1 and trPASS2 are defined:

```
<?xml version="1.0" encoding="UTF-8"?>
<!DOCTYPE tracers SYSTEM "tracers.dtd">
<tracers>
  <passive id="trPASS1">
    <transport type="char">stdaero</transport>
    <unit type="char">kg kg-1</unit>
  </passive>
  <passive id="trPASS2">
    <transport type="char">stdaero</transport>
    <unit type="char">kg kg-1</unit>
  </passive>
</tracers>
```

Additionally, each tracer gets two different meta data. Firstly, transport defines a template of horizontal and vertical advection schemes and flux limiters, in this example the template stdaero is used. A description of available transport templates is given below. Secondly, a unit is specified, which will also be added as meta data to any output of the tracer. Note, that the type of meta data has to be specified. In this example, both meta data are of type character ("char"). For other meta data you could also choose integer ("int") or real ("real").

**Passive Tracers**

For passive tracers, the only required meta data is unit (char). You can find an example for the passive tracer XML file at the runctrl_examples/xml_ctrl folder named tracers_passive.xml.

**Chemical Tracers**

For chemical tracers, the only required meta data is unit (char). Usually, mol_weight (molecular weight, real) is also needed, although it is technically not required. You can find an example for the chemical tracer XML file at the runctrl_examples/xml_ctrl folder named tracers_chemtracer.xml.
Aerosol Tracers

For aerosol tracers, there is a list of necessary meta data specifications: the meta data unit (char), moment (int), mode (char), sol (solubility, real), rho (density, real) and mol_weight (molecular weight, real) are required. You can find an example for the aerosol tracer XML file at the runctrl_examples/xml_ctrl folder named tracers_aerosol.xml.

Available Transport Templates

Currently, there are three different transport templates available: off, stdaero and stdchem. These templates avoid the necessity to add a tracer advection scheme and flux limiter for each single tracer in the namelist. Hence, the values of the namelist parameters ihadv_tracer, ivadv_tracer, itype_hlimit and itype_vlimit are overwritten by the template. Specific information concerning the advection schemes mentioned in the following can be found in Section 3.6.

Specifying off, all advective transport for this tracer is turned off (i.e. ihadv_tracer and ivadv_tracer are set to 0).

The transport template stdaero uses a combination of Miura and Miura with subcycling for the horizontal advection (i.e. ihadv_tracer = 22), 3rd order piecewise parabolic method handling CFL >1 for vertical advection (i.e. ivadv_tracer = 3) in combination with monotonous flux limiters (i.e. itype_hlimit = 4 and itype_vlimit = 4). This means that the conservation of linear correlations is guaranteed which is important for modal aerosol with prognostic mass and number and diagnostic diameter. By this, the diameter of aerosol does not change due to transport.

The transport template stdchem uses the same advection schemes as stdaero (i.e. ihadv_tracer = 22 and ivadv_tracer = 3). However, the considerably faster positive definite flux limiters are used (i.e. itype_hlimit = 3 and itype_vlimit = 2). By this, the mass is still conserved. However, the conservation of linear correlations is traded for a faster computation of the advection.

As you might have noticed in the previous section, the choice of a transport template is not required at the tracer definition. If no transport template is chosen, stdaero is used as default template.

10.4.4. Modes Definition with XML Files

Similar to the previously described tracer definition with XML files, aerosol modes are also defined with XML files. The according namelist parameter specifying the XML file is called cart_modes.xml. An example file modes.xml is provided in the runctrl_examples/xml_ctrl folder. The definition of a mode is done as shown in the following example:

```xml
<?xml version="1.0" encoding="UTF-8"?>
<!DOCTYPE modes SYSTEM "modes.dtd">
```
In this example, a mode called seasa is defined with 2 prognostic moments (2mom). The initial number and mass diameters ($d_{gn}$ and $d_{gm}$) as well as the geometric standard deviation ($\sigma_g$) and density ($\rho$) are specified. For an aerosol tracer, that shall be associated to this mode, the meta data mode has to be set to seasa at the tracer definition (see previous section). In general, all available modes are listed in the example file modes.xml. Hence, it is highly recommended to adapt this file according to your simulation setup.

### 10.4.5. Point Source Module: pntSrc

ICON-ART provides a module which adds emissions from point sources to existing tracers. The namelist switches associated to this module are `lart_passive`, `lart_pntSrc` and `cart_pntSrc.xml` (see Section 10.4.2).

The prerequisite is that you have added a passive tracer via an XML file using the namelist parameter `cart_passive.xml`. Starting from this point, point sources can be added using an XML file specified via `cart_pntSrc.xml`. Additionally, you have to set `lart_passive` and `lart_pntSrc` to .TRUE.. Inside the XML file specified via `cart_pntSrc.xml`, you can add point sources following the subsequent example:

```xml
<?xml version="1.0" encoding="UTF-8"?>
<!DOCTYPE tracers SYSTEM "pntSrc.dtd">

<sources>
    <pntSrc id="RNDFACTORY">
        <substance type="char">testtr</substance>
        <lon type="real">8.00</lon>
        <lat type="real">48.00</lat>
        <source_strength type="real">1.0</source_strength>
        <height type="real">10.</height>
        <unit type="char">kg s^{-1}</unit>
    </pntSrc>
</sources>
```

The options you can specify here have the following meaning:
• **putSrc id**: The name of the point source. This information is actually not used in the ICON-ART code and serves only for a better readability of the XML file. Hence, also multiple point sources with the same id are technically allowed.

• **substance**: This is the name of the substance, the point source emission is added to. Here you have to specify the very same name of the tracer that you have specified in the cart_passive.xml file.

• **lon**: Geographical longitude of the point source in degrees east.

• **lat**: Geographical latitude of the point source in degrees north.

• **source_strength**: The source strength of the point source in the unit specified below.

• **height**: The height of the point source in meters above ground.

• **unit**: The unit of the source strength. Note, that currently every unit different from kg s-1 will lead to a model abort, as no unit conversion is implemented so far.

Via the XML file you can also specify multiple point sources. By this, you can either add point sources to different tracers or specifying multiple source for a single tracer with for example differing source strengths.

### 10.4.6. Volcanic Ash Control

If volcanic eruptions should be considered the switch iart_volcano has to be set. For the 1-moment description of volcanic ash, i.e. six monodisperse size bins for the number concentrations, the integer value is 1. For the 2-moment description where 3 lognormal modes are used, the switch has to be set to 2.

Further input is necessary to define the appropriate volcano(s):

• **volcano_list, whatyouwant.txt** has to contain information for the initialization of each volcano. The path including filename has to be set in the namelist (cart_volcano_file). The line(s) for the volcano(s) of interest can be copied from the file of2009-1133_table3_EDITED_HISTORICAL.txt in the runctrl_examples/emiss_ctrl folder. In Figure 10.1 an example is given. The first column contains an ID of the volcano which is not read but needed due to format specifications. The next column contains the name of the volcano. It is followed by information on its location. There the correct longitude and latitude must be present. Afterwards information on the volcano type is given.

**Figure 10.1.**: Example for the content of volcano_list_name.txt.

With this setup ICON-ART performs a simulation with the standard parameters for the respective volcano type.
10.5. Output

In principle, output of ICON-ART variables works the same way as for ICON variables. The following five quantities of the output have to be specified:

- The time interval between two model outputs.
- The name of the output file.
- The name of the variable(s) and/or variable group(s).
- The type of vertical output grid.
- The type of horizontal output grid.

In general, the output of all (prognostic) tracers defined in the different XML files (passive, chemistry, aerosol) is possible. Additionally, several diagnostic output variables have been added in ICON-ART. These are listed in table 10.6.

There is an option to obtain all variables belonging to a certain group without having to specifying all of them. The output variables that are associated to this group will be written. Available output groups are: ART_AEROSOL, ART_CHEMISTRY, ART_PASSIVE and ART_DIAGNOSTICS. As the names indicate, these groups contain the tracer variables listed in the aerosol-, the chemistry- and the passive-xml, as well as diagnostic variables.

**Table 10.6.:** Selected list of available diagnostic output fields for aerosol.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Associated namelist switch</th>
<th>Description</th>
<th>Groups</th>
</tr>
</thead>
<tbody>
<tr>
<td>seasa_diam</td>
<td>iart_seasalt = 1</td>
<td>Median diameter of sea salt mode A, B, C respectively</td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>seasb_diam</td>
<td>lart_diag_out = .true.</td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>seasc_diam</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>asha_diam</td>
<td>iart_volcano = 2</td>
<td>Median diameter of volcanic ash mode A, B, C respectively</td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>ashb_diam</td>
<td>lart_diag_out = .true.</td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>ashc_diam</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_340nm</td>
<td>iart_volcano = 2</td>
<td>Volcanic ash optical depth at the wavelength indicated by the name</td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_380nm</td>
<td>lart_diag_out = .true.</td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_440nm</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_500nm</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_550nm</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_675nm</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_870nm</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_1020nm</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>tau_volc_1064nm</td>
<td></td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td>ash_total_mc</td>
<td>iart_volcano = 2</td>
<td>Total concentration of volcanic ash in column</td>
<td>ART_DIAGNOSTICS</td>
</tr>
<tr>
<td></td>
<td>lart_diag_out = .true.</td>
<td></td>
<td>ART_DIAGNOSTICS</td>
</tr>
</tbody>
</table>
11. ICON’s Data Assimilation System

In this chapter you will get to know basic components of the ICON data assimilation system. It consists of a whole collection of programs and modules both for the atmospheric variables of the model as well as for soil, snow, ice and sea surface, all collected into the Data Assimilation Coding Environment (DACE).

11.1. Data Assimilation

Numerical weather prediction (NWP) is an initial value problem. The ability to make a skillful forecast heavily depends on an accurate estimate of the present atmospheric state, known as analysis. In general, an analysis is generated by combining, in an optimal way, all available observations with a short term forecast of a general circulation model (e.g. ICON).

Stated in a more abstract way, the basic idea of data assimilation is to fit model states $x$ to observations $y$. Usually, we do not observe model quantities directly or not at the model grid points. Here, we work with observation operators $H$ which take a model state and calculate a simulated observation $y = H(x)$. In terms of software, these model operators can be seen as particular modules, which operate on the ICON model states. Their output is usually written into so-called feedback files, which contain both the real observation $y_{\text{meas}}$ with all its meta data (descriptions, positioning, further information) as well as the simulated observation $y = H(x)$.

However, data assimilation cannot be treated at one point in time only. The information passed on from the past is a crucial ingredient for any data assimilation scheme. Thus, cycling is an important part of data assimilation. It means that we

1. Carry out the core data assimilation component to calculate the so-called analysis $x^{(a)}$, i.e. a state which best fits previous information and the observations $y$,

2. Propagate the analysis $x^{(a)}_{k+1}$ to the next analysis time $t_{k+1}$. Here, it is called first guess or background $x^{(b)}_{k+1}$.

3. Carry out the next analysis by running the core data assimilation component, generating $x^{(a)}_{k+1}$, then cycling the steps.

See Figure 11.1 for a schematic of the basic assimilation process.

11.1.1. Variational Data Assimilation

The basic 3D-VAR step minimizes the functional

$$\mu(x) := \|x - x^{(b)}\|^2_{\mathcal{B}} + \|y - H(x)\|^2_{\mathcal{R}},$$

(11.1)
where $B$ is the background state distribution covariance matrix which is making sure that the information which is available at some place is distributed into its neighborhood properly, and $R$ is the error covariance matrix describing the error distribution for the observations. The minimizer of (11.1) is given by

$$x^{(a)} = x^{(b)} + B H^T (R + B H H^T)^{-1} (y - H(x^{(b)})),$$  \hspace{1cm} (11.2)

The background or first guess $x^{(b)}$ is calculated from earlier analysis by propagating the model from a state $x_{k-1}$ at a previous analysis time $t_{k-1}$ to the current analysis time $t_k$. In the data assimilation code, the minimization of (11.1) is not carried out explicitly by (11.2), but by a conjugate gradient minimization scheme, i.e. in an iterative manner, first solving the equation

$$(R + B H H^T) z_k = y - H(x_k^{(b)}),$$

in observation space calculating $z_k$ at time $t_k$, then projecting the solution back into model space by

$$\delta x_k = x_k^{(a)} - x_k^{(b)} = B H^T z_k.$$ 

We call $\delta x_k$ the analysis increment.

The background covariance matrix $B$ is calculated from previous model runs by statistical methods. We employ the so-called NMC method initially developed by the US weather...
bureau. The matrix $B$ thus contains statistical information about the relationship between different variables of the model, which is used in each of the assimilation steps.

### 11.1.2. Ensemble Kalman Filter

To obtain a better distribution of the information given by observations, modern data assimilation algorithms employ a dynamical estimator for the covariance matrix ($B$-matrix). Given an ensemble of states $x^{(1)}, \ldots, x^{(L)}$, the standard stochastic covariance estimator calculates an estimate for the $B$-matrix by

$$B = \frac{1}{L-1} \sum_{\ell=1}^{L} (x^{(\ell)}_k - \overline{x}_k)(x^{(\ell)}_k - \overline{x}_k)^T,$$

where $\overline{x}$ denotes the mean defined by

$$\overline{x}_k = \frac{1}{L} \sum_{\ell=1}^{L} x^{(\ell)}_k, \quad k \in \mathbb{N}.$$

This is leading us to the Ensemble Kalman Filter (EnKF), where an ensemble is employed for data assimilation and the covariance is estimated by (11.3). Here, we use the name EnKF (ensemble Kalman filter) as a generic name for all methods based on the above idea.

In principle, the EnKF carries out cycling as introduced above, just that the propagation step carries out propagation of a whole ensemble of $L$ atmospheric states $x^{(a,\ell)}_k$ from time $t_k$ to time $t_{k+1}$, and the analysis step has to generate $L$ new analysis members, called the analysis ensemble based on the first guess or background ensemble $x^{(b,\ell)}_k$, $\ell = 1, \ldots, L$.

Usually, the analysis is carried out in observation space, where a transformation is carried out. Also, working with a low number of ensemble members as it is necessary for large-scale data assimilation problems, we need to suppress spurious correlations which arise from a naive application of Eq. (11.3). This technique is known as localization, and the combined transform and localization method is called localized ensemble transform Kalman filter (LETKF), first suggested by Hunt et al. (2007).

The DWD data assimilation coding environment (DACE) provides a state-of-the-art implementation of the LETKF which is equipped with several important ingredients such as different types of covariance inflation. These are needed to properly take care of the modeling error. The original Kalman filter itself does not know what error the model has and thus by default under-estimates this error, which is counter-acted by a collection of tools.

### 11.1.3. Hybrid Data Assimilation

The combination of variational and ensemble methods provides many possibilities to further improve the state estimation of data assimilation. Based on the ensemble Kalman filter LETKF the data assimilation coding environment provides a hybrid system EnVar, the ensemble variational data assimilation.
The basic idea of EnVar is to use the dynamical flow dependent ensemble covariance matrix $B$ as a part of the three-dimensional variational assimilation. Here, localization is a crucial issue, since in the LETKF we localize in observation space, but 3D-VAR employs $B$ in state space. Localization is carried out by a diffusion-type approximation in DACE.

The cycling for the EnVar needs to cycle both the ensemble $x^{(\ell)}$, $\ell = 1, \ldots, L$ and one deterministic state $x_{det}$. The resolution of the ensemble can be lower than the full deterministic resolution. By default we currently employ a 40 km grid spacing for the ensemble and a 13 km global grid spacing for the deterministic state. The ensemble $B$ matrix is then carried over to the finer deterministic resolution by interpolation. See Section 11.2 for more details on the operational assimilation system at DWD.

11.1.4. Surface Analysis

DACE provides additional modules for Sea Surface Temperature (SST) analysis, Soil Moisture Analysis (SMA) and snow analysis. Characteristic time scales of surface and soil processes are typically larger than those of atmospheric processes. Therefore, it is often sufficient to carry out surface analysis only every 6 to 24 hours.

11.2. Assimilation Cycle at DWD

The Assimilation cycle iterates the steps described in Section 11.1: updating a short-range ICON forecast (first guess) using the observations available for that time window to generate an analysis, from which then a new updated first guess is started.

The core assimilation for atmospheric fields is based on a hybrid system (EnVar) as described in Section 11.1.3. At every assimilation step (every 3 h) an LETKF is ran using an ensemble of ICON first guesses. Currently, the ensemble consists of 40 members with a horizontal grid spacing of 40 km and a 20 km nest over Europe. A convex linear combination of the 3D-VAR climatological and the LETKF’s (flow dependent) covariance matrix is then used to run a deterministic 3D-VAR analysis at 13 km horizontal grid spacing.

In addition, the above mentioned surface modules are ran: Sea Surface Temperature (SST) analysis, Soil Moisture Analysis (SMA) and snow analysis.

Note that for the ICON-EU nest no assimilation of atmospheric fields is conducted. Instead the necessary atmospheric analysis increments are interpolated from the underlying global grid. Together with the available first guess fields on the nest they form the nest analysis. A separate surface analysis, however, is conducted.

The deterministic as well as the ensemble analysis is generated 8 times a day. Based on the former, deterministic forecasts are launched at approx. 13 km horizontal grid spacing globally with a 6.5 km nest over Europe. The maximum forecast time of the whole system is limited to +30 h lead time at 03/09/15/21 UTC. Otherwise, the system is integrated up to +120 h while at 00/12 UTC the integration on the global domain (only) is prolonged to +180 h lead time.
Since the beginning of 2018 ICON ensemble forecasts are conducted as well. On the basis of the analysis ensemble 40 short to medium forecasts at 40 km globally and 20 km nested over Europe are run 8 times a day. The maximum forecast times are equivalent to those of the deterministic system (see above). The primary purpose of the ensemble forecasts is to estimate the forecast uncertainty, which arises due to uncertainties in the initial conditions and the model error.

The input, output and processes involved in the assimilation cycle are briefly described below:

**Atmospheric Analysis**

**Fields modified by the atmospheric analysis:** (see Appendix B for a description of each variable) t, p, u, v, qv.

**Grid(s) on which it is performed:** global

Carried out at every assimilation time step (3 h) using the data assimilation algorithms described in the previous sections.

**Main input:** First guess, observations, previous analysis error, online bias correction files.

**Main output:** Analysis of the atmospheric fields, analysis error, bias correction files, feedback files with information on the observation, its departures to first guess and analysis.

The system can make use of the following observations: radiosondes, weather stations, buoys, aircraft, ships, radio occultations, AMV winds and radiances. Available general features of the module are variational quality control and (variational) online bias correction. Regarding EnKF specifics, different types of inflation techniques, relaxation to prior perturbations and spread, adaptive localization, SST perturbations and SMA perturbations are available.

**Snow Analysis**

**Fields modified by the snow analysis:** (see Appendix B for a description of each variable) freshsnow, h_snow, rho_snow, t_snow, w_i, w_snow.

**Grid(s) on which it is performed:** global, EU-nest

Carried out at each assimilation time step (3 h).

**Main input:** SYNOP snow depth observations if the coverage is sufficient. If this is not the case, more sources of information are looked for until the number of observations is high enough, namely (and in this order), precipitation and 2 m temperature, direct observations (wwreports) and the NCEP external snow analysis.

**Main output:** Analysis of the snow fields.
**Sea Surface Temperature Analysis**

**Fields modified by the SST analysis:** (see Appendix B for a description of each variable) \( fr_{\text{seaice}}, h_{\text{ice}}, t_{\text{ice}}, t_{\text{so}} \).

**Grid(s) on which it is performed:** global, EU-nest

Carried out only once a day, at 0 UTC.

**Main input:** NCEP analysis from the previous day (which uses satellite, buoy and ship observations, to be used as a first guess), ship and buoy observations available since the time of the NCEP analysis.

**Main output:** Sea surface temperature analysis and estimated error.

**Soil Moisture Analysis**

**Fields modified by the SMA analysis:** (see Appendix B for a description of each variable) \( w_{\text{so}} \).

**Grid(s) on which it is performed:** global, EU-nest

Carried out only once a day, at 0 UTC.

**Main input:** Background fields for relevant fields at every hour since last assimilation, 2 m-temperature analysis (see below) to be used as observations.

**Main output:** Soil moisture analysis and estimated error.

**2m-Temperature Analysis**

Although carried out only at 0 UTC, it is run for several time steps in between to provide the output (2 m temperature) needed by the SMA analysis. Uses observations from SYNOP stations on land and METAR information from airports.
A. The Computer System at DWD

Available Platforms at DWD

The NWP exercises will be mainly carried out on the Cray XC 40 supercomputer system at DWD. This system consists of several compute nodes with corresponding service nodes.

- **xce00.dwd.de, xce01.dwd.de:**
  These are the login nodes, which run a SUSE Linux Enterprise (SLES) Linux. The nodes are used for **compiling and linking**, preparation of data and basic editing work. They are not used for running parallel programs, but jobs can be submitted to the Cray XC 40 compute nodes.

- **Cray XC 40:**
  The Cray XC 40 has 432 **compute nodes**, where each node is equipped with 2 Intel Haswell processors with 12 cores and a second node partition with 864 Intel Broadwell nodes. Each Haswell node therefore has 24 computational cores (Broadwell: 36). These nodes cannot be accessed interactively, but only by **batch jobs**. Such jobs can use up to 62 GByte of main memory per node, which is about 2.5 GByte per core (Broadwell: 1.7 GByte). For normal test jobs it should be enough to use 10-15 nodes (depending on the chosen grid resolution).

- **lce00.dwd.de, lce01.dwd.de:**
  Tools for the **visualization of meteorological fields** (CDO, NCL, ncview) are only available on the pre-/post-processing cluster lce. This cluster comprises 66 Haswell batch nodes, but we will only use the login nodes lce00, lce01 during the course.

There is a common filesystem across all nodes and every user has three different main directories:

- **/e/uhome/username ($HOME)**
  Directory for storing source code and scripts to run the model. This is a Panasas filesystem suitable for many small files.

- **/e/uwork/username ($WORK)**
  Directory for storing larger amounts of data.

- **/e/uscratch/username ($SCRATCH)**
  Directory for storing very large amounts of data. For the $WORK and the $SCRATCH filesystem there is a common quota for every user of 2.5 TByte.
The Batch System for the Cray XC 40

Jobs for the Cray XC 40 system have to be submitted with the batch system PBS. These batch jobs may either be launched from the Linux cluster lce or from the XC 40 login nodes xce00/01. Together with the source code of the programs we provide some run scripts in which all necessary batch commands are set.

Here are the most important commands for working with the PBS:

- **qsub job_name** to submit a batch job to PBS. This is done in the run scripts.
- **qstat** to query the status of all batch jobs on the XC 40. You can see whether jobs are Q (queued) or R (running). You have to submit jobs to the queue xc_norm.h.
- **qstat -u user** to query the status of all your batch jobs on the machine.
- **qdel job_nr@machine** to cancel your job(s) from the batch queue of a machine. The job_nr is given by qstat -w.
- **qcat -f job_nr@machine** To follow stdout and stderr interactively.

In your run scripts, execution begins in your home directory, regardless of what directory your script resides in or where you submitted the job from. You can use the **cd** command to change to a different directory. The environment variable **$PBS_O_WORKDIR** makes it easy to return to the directory from which you submitted the job:

```
cd $PBS_O_WORKDIR
```
### B. Table of ICON Output Variables

The following table contains the NWP variables available for output\(^1\). Please note that the field names are following an ICON-internal nomenclature, see Section 7.1 for details. The list also contains tile-based fields (suffix \_t1, \_t2, \ldots) which are summarized as \_t*.

By "ICON-internal" variable names, we denote those field names that are provided as the string argument \texttt{name} to the subroutine calls \texttt{CALL add\_var(...) and \texttt{CALL add\_ref(...) inside the ICON source code. These subroutine calls have the purpose to register new variables, to allocate the necessary memory, and to set the meta-data for these variables.}

<table>
<thead>
<tr>
<th>Variable Name</th>
<th>GRIB2 Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>acdnc</td>
<td>ndcloud</td>
<td>Cloud droplet number concentration</td>
</tr>
<tr>
<td>adrag_u_grid</td>
<td>Zonal resolved surface stress mean since model start</td>
<td></td>
</tr>
<tr>
<td>adrag_v_grid</td>
<td>Meridional resolved surface stress mean since model start</td>
<td></td>
</tr>
<tr>
<td>aer_bc</td>
<td>aer_bc</td>
<td>Black carbon aerosol</td>
</tr>
<tr>
<td>aer_du</td>
<td>aer_dust</td>
<td>Total soil dust aerosol</td>
</tr>
<tr>
<td>aer_or</td>
<td>aer_org</td>
<td>Organic aerosol</td>
</tr>
<tr>
<td>aer_ss</td>
<td>aer_ss</td>
<td>Sea salt aerosol</td>
</tr>
<tr>
<td>aer_su</td>
<td>aer_so4</td>
<td>Total sulfate aerosol</td>
</tr>
<tr>
<td>aercl_bc</td>
<td>Black carbon aerosol climatology</td>
<td></td>
</tr>
<tr>
<td>aercl_du</td>
<td>Total soil dust aerosol climatology</td>
<td></td>
</tr>
<tr>
<td>aercl_or</td>
<td>Organic aerosol climatology</td>
<td></td>
</tr>
<tr>
<td>aercl_ss</td>
<td>Sea salt aerosol climatology</td>
<td></td>
</tr>
<tr>
<td>aercl_su</td>
<td>Total sulfate aerosol climatology</td>
<td></td>
</tr>
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<td>albdif</td>
<td>alb_dif</td>
<td>Shortwave albedo for diffuse radiation</td>
</tr>
<tr>
<td>albsi</td>
<td>alb_seaice</td>
<td>Sea ice albedo (diffuse)</td>
</tr>
<tr>
<td>albdif_t*</td>
<td>alb_rad</td>
<td>Tile-based shortwave albedo for diffusive radiation</td>
</tr>
<tr>
<td>albni</td>
<td>alb_ni</td>
<td>Near IR albedo for diffuse radiation</td>
</tr>
<tr>
<td>albnirdif_t*</td>
<td>alb_ni</td>
<td>Tile-based near IR albedo for diffuse radiation</td>
</tr>
<tr>
<td>albnirdir</td>
<td>aizin</td>
<td>Near IR albedo for direct radiation</td>
</tr>
<tr>
<td>albuuv</td>
<td>alb_uv</td>
<td>UV visible albedo for diffuse radiation</td>
</tr>
<tr>
<td>albvisdif_t*</td>
<td>alb_uv</td>
<td>Tile-based UV visible albedo for diffusive radiation</td>
</tr>
<tr>
<td>albvisdif</td>
<td>alb_uv</td>
<td>UV visible albedo for diffuse radiation</td>
</tr>
<tr>
<td>albvisdir</td>
<td>aluvp</td>
<td>UV visible albedo for direct radiation</td>
</tr>
<tr>
<td>alhfl_b</td>
<td>alhfl_b</td>
<td>Latent heat flux from bare mean since model start</td>
</tr>
<tr>
<td>alhfl_p</td>
<td>alhfl_p</td>
<td>Latent heat flux from plantmean since model start</td>
</tr>
<tr>
<td>alhfl_s</td>
<td>alhfl_s</td>
<td>surface latent heat flux mean since model start</td>
</tr>
<tr>
<td>aqhfl_s</td>
<td>evapt</td>
<td>surface moisture flux mean since model start</td>
</tr>
</tbody>
</table>

---

\(^1\)The Table B.1 in this Appendix is based on the revision state 77cdb449 (2019-02-27).
### Table B.1 – Continued from previous page

<table>
<thead>
<tr>
<th>Variable Name</th>
<th>GRIB2 Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ashfl_s</td>
<td>ashfl_s</td>
<td>Surface sensible heat flux mean since model start</td>
</tr>
<tr>
<td>asob_s</td>
<td>asob_s</td>
<td>Surface net solar radiation mean since model start</td>
</tr>
<tr>
<td>asob_t</td>
<td>asob_t</td>
<td>TOA net solar radiation mean since model start</td>
</tr>
<tr>
<td>asod_t</td>
<td>sodt_rad</td>
<td>Top down solar radiation mean since model start</td>
</tr>
<tr>
<td>asodifd_s</td>
<td>aswdifd_s</td>
<td>Surface down solar diff. rad. mean since model start</td>
</tr>
<tr>
<td>asodifu_s</td>
<td>aswdifu_s</td>
<td>Surface up solar diff. rad. mean since model start</td>
</tr>
<tr>
<td>asodird_s</td>
<td>aswdir_s</td>
<td>Surface down solar direct rad. mean since model start</td>
</tr>
<tr>
<td>asou_t</td>
<td>usurf</td>
<td>Top up solar radiation mean since model start</td>
</tr>
<tr>
<td>astr_u_sso</td>
<td>lgvs</td>
<td>Zonal sso surface stress mean since model start</td>
</tr>
<tr>
<td>astr_v_sso</td>
<td>mgvs</td>
<td>Meridional sso surface stress mean since model start</td>
</tr>
<tr>
<td>aswflx_par_sfc</td>
<td>apab_s</td>
<td>Downward PAR flux mean since model start</td>
</tr>
<tr>
<td>athb_s</td>
<td>athb_s</td>
<td>Surface net thermal radiation mean since model start</td>
</tr>
<tr>
<td>athb_t</td>
<td>athb_t</td>
<td>TOA net thermal radiation mean since model start</td>
</tr>
<tr>
<td>athd_s</td>
<td>athd_s</td>
<td>Surface down thermal radiation mean since model start</td>
</tr>
<tr>
<td>athu_s</td>
<td>athu_s</td>
<td>Surface up thermal radiation mean since model start</td>
</tr>
<tr>
<td>asmfl_s</td>
<td>asmfl_s</td>
<td>U-momentum flux flux at sumean since model start</td>
</tr>
<tr>
<td>avg_qc</td>
<td>tqc_dia</td>
<td>Tci specific cloud water content (diagnostic) avg</td>
</tr>
<tr>
<td>avg_qi</td>
<td>tqi_dia</td>
<td>Tci specific cloud ice content (diagnostic) avg</td>
</tr>
<tr>
<td>avg_qv</td>
<td>tqv_dia</td>
<td>Column integrated water vapour (diagnostic) avg</td>
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<td>asmfl_s</td>
<td>asmfl_s</td>
<td>V-momentum flux flux at sumean since model start</td>
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<td>c_t_lk</td>
<td>Shape factor (temp. profile in lake thermocline)</td>
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<td>cape_ml</td>
<td>cape_ml</td>
<td>Cape of mean surface layer parcel</td>
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<td>cape</td>
<td>cape_con</td>
<td>Conv avail pot energy</td>
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<td>Convective inhibition of mean surface layer parcel</td>
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<td>Low level clouds</td>
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<td>clicm</td>
<td>Mid level clouds</td>
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<td>Total cloud cover time avg</td>
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<td>Modified total cloud cover for media</td>
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<td>clct</td>
<td>Total cloud cover</td>
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<td>clic</td>
<td>Cloud cover</td>
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<td>cldepth</td>
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<td>cloud_num</td>
<td>Cloud droplet number concentration</td>
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<td>conv Gust</td>
<td>conv Gust</td>
<td>Convective contribution to wind gust</td>
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<tr>
<td>cosmu0</td>
<td>uvcossza</td>
<td>Cosine of solar zenith angle</td>
</tr>
<tr>
<td>ddt_adv_q</td>
<td>ddt_adv_q</td>
<td>Adveective tracer tendency</td>
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<tr>
<td>ddt_exner_phy</td>
<td>ddt_exner_phy</td>
<td>Exner pressure physical tendency</td>
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<td>ddt_grf_q</td>
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<td>Tracer tendency for grid refinement</td>
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<td>ddt_pres_sfc</td>
<td>Surface pressure tendency</td>
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<td>ddt_qc_conv</td>
<td>Convective tendency of cloud water mass density</td>
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<td>ddt_qc_gscp</td>
<td>Microphysics tendency of specific cloud water</td>
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<td>Turbulence tendency of specific cloud water</td>
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<tr>
<td>ddt_qi_conv</td>
<td>ddt_qi_conv</td>
<td>Convective tendency of cloud ice mass density</td>
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<tr>
<td>ddt_qi_gscp</td>
<td>ddt_qi_gscp</td>
<td>Microphysics tendency of specific cloud ice</td>
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<td>ddt_qi_turb</td>
<td>ddt_qi_turb</td>
<td>Turbulence tendency of specific cloud ice</td>
</tr>
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<td>Convective tendency of rain mass density</td>
</tr>
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<td>ddt_nr_gscp</td>
<td>Microphysics tendency of rain</td>
</tr>
<tr>
<td>Variable Name</td>
<td>GRIB2 Name</td>
<td>Description</td>
</tr>
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</tr>
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<td>Convective tendency of snow mass density</td>
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<td>Microphysics tendency of snow</td>
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<td>Microphysics tendency of specific humidity</td>
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<td>ttends</td>
<td>Sso + gwdrag temperature tendency</td>
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<td>Dynamical temperature tendency</td>
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<td>Microphysical temperature tendency</td>
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<td>Long wave radiative temperature tendency</td>
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<td>Short wave radiative temperature tendency</td>
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<td>Geopotential difference between half levels</td>
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<td>div_ic</td>
<td>rdiv</td>
<td>Divergence at half levels</td>
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<td></td>
<td>Divergence</td>
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<td></td>
<td>Depth of thermally active layer of bot. sediments.</td>
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<td></td>
<td>Pressure thickness</td>
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<td>drag_u_grid</td>
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<td>Meridional resolved surface stress</td>
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<td>dtheta_v_ic_ubc</td>
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<td>Potential temperature at child upper boundary</td>
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<tr>
<td>dvn-ie_int</td>
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<td>Normal velocity at parent interface level</td>
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<tr>
<td>dvn-ie_ubc</td>
<td></td>
<td>Normal velocity at child upper boundary</td>
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<td>du_ubc</td>
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<td>Vertical velocity at child upper boundary</td>
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<td>Zonal gradient of vertical wind</td>
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<tr>
<td>dudy</td>
<td></td>
<td>Meridional gradient of vertical wind</td>
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<td>dyn_gust</td>
<td></td>
<td>Dynamical gust</td>
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Continued on next page
### Variable Name | GRIB2 Name | Description
--- | --- | ---
fetch lk | fetch lk | Wind fetch over lake
fis | fis | Geopotential (s)
for d | for_d | Fraction of deciduous forest
fr glac | fr_glac | Fraction glacier
fr lake | fr_lake | Fraction lake
fr land | fr_land | Fraction land
fr seaice | fr_ice | Fraction of sea ice
frac t_* | fr_luc | Tile point area fraction list
freshsnow t_* | freshsnow | Indicator for age of snow in top of snow layer
freshsnow | freshsnow | Weighted indicator for age of snow in top of snow layer
gamso lk |  | Attenuation coefficient of lake water with respect to sol. rad.
geopot agl ifc | fi | Geopotential above groundlevel at cell center
geopot agl | fi | Geopotential above groundlevel at cell center
geopot | fi | Geopotential at full level cell centre
graupel gsp rate | prg_gsp | Gridscale graupel rate
grau gsp | grau_gsp | Gridscale graupel
grf tend mfix | graftmfix | Normal mass flux tendency (grid refinement)
grf tend rho | graftrho | Density tendency (grid refinement)
grf tend thv | graftthv | Virtual potential temperature tendency (grid refinement)
grf tend vn | graftvn | Normal wind tendency (grid refinement)
grf tend w | gravity | Vertical wind tendency (grid refinement)
gust | vmax_10m | Gust at 10 m
gz0 t_* | z0 | Tile-based roughness length times gravity
gz0 | z0 | Roughness length
h bli lk | h_bli lk | Thickness of the upper layer of the sediments
h ice | h_ice | Sea/lake-ice depth
h ml lk | h_mli lk | Mixed-layer thickness
h snow lk | h_snowlk | Depth of snow on lake ice
h snow si | h_snowsi | Depth of snow on sea ice
h snow t_* | h_snow | Snow height
h snow | h_snow | Weighted snow depth
hbas con | hbas_con | Height of convective cloud base
hdef ic | deformation | Deformation
hf1 q | horizontal | Horizontal tracer flux
hno3 | hno3 | Height of O3 maximum (Pa)
htop con | htop_con | Height of convective cloud top
htop dc | htop_dc | Height of top of dry convection
hzero cl | hzerocl | Height of 0 deg C level
k400 |  | Level index corresponding to the HAG of the 400hPa level
k800 |  | Level index corresponding to the HAG of the 800hPa level
k850 |  | Level index corresponding to the HAG of the 850hPa level
k950 |  | Level index corresponding to the HAG of the 950hPa level

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<td>tkvm</td>
<td>Turbulant diffusion coefficients for momentum</td>
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<td>tmax_2m</td>
<td>Max 2m temperature</td>
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<tr>
<td>tmin_2m</td>
<td>tmin_2m</td>
<td>Min 2m temperature</td>
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<tr>
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<td>Geometric height of the earths surface above sea level</td>
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<td>tot_pr</td>
<td>Total precip rate, time average</td>
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<td>tot_prec</td>
<td>Total precip</td>
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<td>qc_dia</td>
<td>Total specific cloud water content (diagnostic)</td>
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<tr>
<td>tot_qi_dia</td>
<td>qi_dia</td>
<td>Total specific cloud ice content (diagnostic)</td>
</tr>
<tr>
<td>tot_qv_dia</td>
<td>qv_dia</td>
<td>Total specific humidity (diagnostic)</td>
</tr>
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<td>tqc</td>
<td>tqc</td>
<td>Total column integrated cloud water</td>
</tr>
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<td>Total column integrated graupel</td>
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<td>tqi_dia</td>
<td>Total column integrated cloud ice (diagnostic)</td>
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<td>tqi</td>
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<td>tqv_dia</td>
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<td>Total column integrated water vapour</td>
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<td>Average of vertically integrated tracers</td>
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<td>trsolall</td>
<td></td>
<td>Shortwave net transmissivity</td>
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<td>Reference surface temperature</td>
</tr>
<tr>
<td>tsfctrad</td>
<td></td>
<td>Surface temperature at trad</td>
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<tr>
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<td>Tile-based turbulent transfer velocity for heat</td>
</tr>
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<td>tvh</td>
<td></td>
<td>Turbulent transfer velocity for heat</td>
</tr>
<tr>
<td>tvvm_t_</td>
<td></td>
<td>Tile-based turbulent transfer velocity for momentum</td>
</tr>
<tr>
<td>tvm</td>
<td></td>
<td>Turbulent transfer velocity for momentum</td>
</tr>
<tr>
<td>u_10m_t_</td>
<td>u_10m</td>
<td>Tile-based zonal wind in 2m</td>
</tr>
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<td>u_10m</td>
<td>u_10m</td>
<td>Zonal wind in 10m</td>
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<tr>
<td>umfl_s_t_</td>
<td>umfl_s</td>
<td>U-momentum flux at the surface</td>
</tr>
<tr>
<td>umfl_s</td>
<td>umfl_s</td>
<td>U-momentum flux at the surface</td>
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<tr>
<td>u</td>
<td>u</td>
<td>Zonal wind</td>
</tr>
<tr>
<td>v_10m_t_</td>
<td>v_10m</td>
<td>Tile-based meridional wind in 2m</td>
</tr>
<tr>
<td>v_10m</td>
<td>v_10m</td>
<td>Meridional wind in 10m</td>
</tr>
<tr>
<td>vfl_q</td>
<td></td>
<td>Vertical tracer flux</td>
</tr>
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<table>
<thead>
<tr>
<th>Variable Name</th>
<th>GRIB2 Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>vio3</td>
<td>tozne</td>
<td>Vertically integrated ozone amount</td>
</tr>
<tr>
<td>vmfl_s_t_*</td>
<td>vmfl_s</td>
<td>V-momentum flux at the surface</td>
</tr>
<tr>
<td>vn_ie</td>
<td>vn</td>
<td>Normal wind at half level</td>
</tr>
<tr>
<td>vn</td>
<td>relv</td>
<td>Velocity normal to edge</td>
</tr>
<tr>
<td>vor</td>
<td>relv</td>
<td>Vorticity</td>
</tr>
<tr>
<td>vt</td>
<td>vt</td>
<td>Tangential-component of wind</td>
</tr>
<tr>
<td>vwind_expl_wgt</td>
<td>v</td>
<td>Explicit weight in vertical wind solver</td>
</tr>
<tr>
<td>vwind_impl_wgt</td>
<td>v</td>
<td>Implicit weight in vertical wind solver</td>
</tr>
<tr>
<td>v</td>
<td>v</td>
<td>Meridional wind</td>
</tr>
<tr>
<td>w_concorr_c</td>
<td>w_i</td>
<td>Contravariant vertical correction</td>
</tr>
<tr>
<td>w_i_t_*</td>
<td>w_i</td>
<td>Weighted water content of interception water</td>
</tr>
<tr>
<td>w_i</td>
<td>w_i</td>
<td>Weighted water content of interception water</td>
</tr>
<tr>
<td>w_snow_t_*</td>
<td>w_snow</td>
<td>Water equivalent of snow</td>
</tr>
<tr>
<td>w_snow</td>
<td>w_snow</td>
<td>Weighted water equivalent of snow</td>
</tr>
<tr>
<td>w_so_ice_t_*</td>
<td>w_so_ice</td>
<td>Ice content</td>
</tr>
<tr>
<td>w_so_ice</td>
<td>w_so_ice</td>
<td>Ice content</td>
</tr>
<tr>
<td>w_so_t_*</td>
<td>w_so</td>
<td>Total water content (ice + liquid water)</td>
</tr>
<tr>
<td>w_so</td>
<td>w_so</td>
<td>Total water content (ice + liquid water)</td>
</tr>
<tr>
<td>ww</td>
<td>ww</td>
<td>Significant weather</td>
</tr>
<tr>
<td>w</td>
<td>w</td>
<td>Vertical velocity</td>
</tr>
<tr>
<td>z_ifc</td>
<td>hhl</td>
<td>Geometric height at half level center</td>
</tr>
<tr>
<td>z_mc</td>
<td>h</td>
<td>Geometric height at full level center</td>
</tr>
</tbody>
</table>

### List of ICON Variable Groups

The "group:" keyword for the namelist parameters `ml_varlist, hl_varlist, pl_varlist` (namelist `output_nml`) can be used to activate a set of common variables for output at once. The following lists contain the variables for each of these groups (empty groups and groups with only a single entry are omitted).

**Group ADDITIONAL_PRECIP_VARS**
- cape, clct, clct_mod, prec_con_rate_avg, prec_gsp_rate_avg, tqc_dia, tqi_dia, tqv_dia

**Group ATMO_DERIVED_VARS**
- div, omega_z, vor

**Group ATMO_ML_VARS**
- pres, qc, qg, qi, qr, qs, qv, temp, tke, u, v, w

**Group ATMO_PL_VARS**
- qc, qg, qi, qr, qs, qv, temp, tke, u, v, w

**Group ATMO_ZL_VARS**
- pres, qc, qg, qi, qr, qs, qv, temp, tke, u, v, w

**Group CLOUD_DIAG**
- clc, tot_qc_dia, tot_qi_dia, tot_qv_dia
Group DWD_FG_ATM_VARS
  pres, pres_sfc, qc, qi, qr, qs, qv, rho, t_2m, td_2m, temp, theta_v, tke, u, u_10m, v, v_10m, vn, w, z_ifc

Group DWD_FG_SFC_VARS
  alb_si, c_t_lk, fr_land, fr_seaice, freshsnow, gz0, h_ice, h_ml lk, h_snow, hsnow_max, plantevap, qv_s, rho_snow, snow_age, snowfrac_lc, t_bot lk, t_g, t_ice, t_mnw lk, t_seasfc, t_snow, t_so, t_wml lk, w_i, w_snow, w_so, w_so_ice

Group DWD_FG_SFC_VARS_T
  freshsnow_t_*, h_snow_t_*, plantevap_t_*, qv_s_t_*, rho_snow_t_*, snowfrac_lc_t_*, t_g_t_*, t_snow_t_*, t_so_t_*, w_i_t_*, w_snow_t_*, w_so_ice_t_*, w_so_t_*

Group ICON_LBC_VARS
  pres, qc, qi, qr, qs, qv, temp, tke, u, v, w, z_ifc

Group LAND_TILE_VARS
  h_snow_t_*, plantevap_t_*, qv_s_t_*, rho_snow_t_*, snowfrac_lc_t_*, snowfrac_t_*, t_g_t_*, t_s_t_*, t_snow_t_*, t_so_t_*, w_i_t_*, w_snow_t_*, w_so_ice_t_*, w_so_t_*

Group LAND_VARS
  qv_s, rho_snow, snowfrac, snowfrac_lc_t, t_g, t_snow, t_so, w_i, w_snow, w_so, w_so_ice

Group LATBC_PREFETCH_VARS
  pres, pres_sfc, qc, qi, qr, qs, qv, rho, temp, theta_v, u, v, vn, w, z_ifc

Group MODE_COMBINED_IN
  fr_seaice, freshsnow, h_ice, h_snow, qv_s, rho_snow, t_g, t_ice, t_snow, t_so, w_i, w_snow, w_so

Group MODE_COSMO_IN
  alb_si, freshsnow, h_ice, qv_s, rho_snow, t_g, t_ice, t_snow, t_so, w_i, w_snow, w_so

Group MODE_DWD_ANA_IN
  fr_seaice, freshsnow, h_ice, h_snow, pres, qv, t_ice, t_seasfc, t_snow, t_so, temp, u, v, w, w_so

Group MODE_DWD_FG_IN
  alb_si, c_t_lk, gz0, h_ml lk, hsnow_max, plantevap, qc, qg, qi, qr, qs, qv, rho, rho_snow, snow_age, snowfrac_lc, t_bot lk, t_g, t_ice, t_mnw lk, t_snow, t_so, t_wml lk, theta_v, tke, vn, w, w_i, w_snow, w_so_ice, z_ifc

Group MODE_IAU_ANAATM_IN
  pres, qc, qi, qv, temp, u, v

Group MODE_IAU_ANA_IN
  fr_seaice, freshsnow, h_snow, pres, qc, qi, qv, t_seasfc, t_so, temp, u, v, w_so

Group MODE_IAU_FG_IN
  alb_si, c_t_lk, freshsnow, gz0, h_ice, h_ml lk, h_snow, hsnow_max, plantevap, qc, qg, qi, qr, qs, qv, rho, rho_snow, snow_age, snowfrac_lc, t_bot lk, t_g, t_ice, t_mnw lk, t_snow, t_so, t_wml lk, theta_v, tke, vn, w, w_i, w_snow, w_so_ice, w_so_ice
B. Output Variables

Group MODE_IAU_OLD_ANA_IN
fr_seaice, freshsnow, h_snow, pres, qv, rho_snow, t_seasfc, t_so, temp, u, v, w_snow, w_so

Group MODE_IAU_OLD_FG_IN
alb_si, c_t lk, gz0, h_ice, h_ml lk, qc, qi, qr, qs, qv, rho, t_bot lk, t_g, t_ice, t_mnw lk, t_snow, t_so, t_wml lk, theta_v, tke, vn, w, w_i, w_so, w_so_ice

Group MODE_INIANA
alb_si, c_t lk, fr_land, fr_seaice, freshsnow, gz0, h_ice, h_ml lk, h_snow, hsnow_max, plantevap, pres, qc, qi, qr, qs, qv, rho_snow, smi, snow_age, t_bot lk, t_g, t_ice, t_mnw lk, t_snow, t_so, t_wml lk, temp, tke, u, v, w, w_i, w_snow, w_so_ice, z_ifc

Group NH_PROG_VARS
exner, rho, theta_v, vn

Group PBL_VARS
alhfl_s, aqhfl_s, ashfl_s, gust10, lhfl bs, lhfl_s, qhfl_s, qv_2m, shfl_s, t_2m, t_2m_land, tch, tcm, td_2m, td_2m_land, tkr, tkvh, tkvm, tvh, tvm, u_10m, v_10m

Group PHYS_TENDENCIES
ddt_temp_drag, ddt_temp_radlw, ddt_temp_radsw, ddt_temp_turb, ddt_u_gwd, ddt_u_sso, ddt_u_turb, ddt_v_gwd, ddt_v_sso, ddt_v_turb

Group PRECIP_VARS
graupel_gsp, prec_con, prec_gsp, rain_con, rain_gsp, snow_con, snow_gsp, tot_prec

Group RAD_VARS
albdif, albnirdif, albvisdif, asob_s, asob_t, asod_t, asodifd_s, asodifu_s, asodird_s, asou_t, aswflx_par_sfc, athb_s, athb_t, athd_s, athu_s, sob_s, sob_t, sod_t, soddifd_s, sou_s, sou_t, swflx_par_sfc, thb_s, thb_s_t_*, thu_s

Group SNOW_VARS
rho_snow, t_snow
B. Output Variables
# C. Exercises

On the following pages a number of exercises is provided. These exercises revisit the topics of the individual chapters and range from easy tests to the setup of complex forecast simulations.

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<td>Ex. C.4.1: Running idealized test cases Ex. C.4.2: Nested sub-domains Ex. C.4.3: Tracer advection</td>
</tr>
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<td>Ex. C.2.1: Grid generator and ExtPar web interface</td>
<td>Ex. C.2.2: Retrieving DWD analysis data Ex. C.2.3: Retrieving IFS data</td>
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<tr>
<td>Real Data Runs</td>
<td>Ex. C.5.1: Global ICON forecast from DWD analysis Ex. C.5.2: Time-stepping Ex. C.5.3: Run-time performance Ex. C.5.4: Parallel scaling Ex. C.5.5: Writing output for driving ICON-LAM Ex. C.5.6: Spin-up effects Ex. C.5.7: Bit-reproducibility</td>
</tr>
<tr>
<td>Ex. C.2.4: Preparation of limited area run</td>
<td></td>
</tr>
<tr>
<td>ICON-LAM</td>
<td>Ex. C.6.1: Limited area run Ex. C.6.2: Modifying the temporal resolution Ex. C.6.3: Driving ICON-LAM by DWD Data</td>
</tr>
<tr>
<td>Programming ICON</td>
<td>Ex. C.8.1: Allocating Additional Fields Ex. C.8.2: Looping over grid points Ex. C.8.3: Implementing own diagnostics</td>
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C.1. Installation of the ICON Model Package

For practical work during these exercises, you need information about the use of the computer systems and from where you can access the necessary files and data. Please take a look at Appendix A to get information on how to use DWD’s supercomputer and run test jobs.

In this exercise we will extract and build the ICON model and its pre-processing tools:

EX 1.1

Log into the Cray XC 40 login node “xce” and install the ICON model in the $WORK directory of your Cray XC 40 user account. For that you have to do the following:

- The necessary files for the ICON tutorial can be found in the subdirectory
  
  /e/uwork/trng024/packages

  Copy the tar-file icon_tutorial.tar.gz into your $WORK directory.

- Change into your $WORK directory and extract the compressed tar file to yield the directory structure depicted in Figure 1.1 containing the ICON sources and test data.

- Follow the instructions in Section 1.2.2 to configure and compile the ICON model on the Cray XC 40 platform.

  Note: Note that you can also compile the routines in parallel by using the GNU-make with the command gmake -j np

Install also the DWD ICON Tools:

- Change into the subdirectory dwd_icon_tools and build the DWD ICON Tools according to Section 1.3.2.

C.2. Necessary Input Data

In the following exercises, you will deal with the necessary preparatory steps for performing global real case ICON runs.

Preparation of Global Runs

In this exercise we will generate the necessary grids and external parameter files using the ICON web service (see Section 2.1.4):
EX 2.1 Grid generator and ExtPar web interface:

- Open the DWD grid generator and ExtPar web interface
  
  https://webservice.dwd.de/cgi-bin/spp1167/webservice.cgi

  in your web browser.

- Select a sequence of three grids: Domain #0 – reduced radiation grid, domain #1 – global grid, domain #2 – refined region over Europe.

  Make sure that you set the “parent grid ID” appropriately.

- For the reduced radiation grid, select a “synthetic grid” with R3B05 resolution.

  Select global refinement for the domain #1.

- Specify the R3B07 grid, domain #2, as a regular-shaped refinement region over Europe, see the illustration on the right or Fig. C.1:

  Center lon/lat: 19.5 / 50.0 deg, half width lon/lat: 44.5 / 21.5 deg.

- Submit the web form – but do not forget to provide your e-mail address! Wait for the job to finish, this may take some time. Afterwards, pick the generated data according to the download information that you have received via e-mail.

- Copy and rename the grid files s. t. the file names match the nomenclature iconR<nroot>B<jlev>_DOM<idom>.nc, see Section 5.1.2.

  Rename the ExtPar data to extpar_DOM<idom>.nc.

- Take a look at the grid data and the external parameters using the ncdump utility, see Section 9.1.1 for details. Find out whether the external parameter fields are defined at the grid vertices, edge midpoints or cells.

In this exercise we will retrieve and pre-process DWD analysis data for global forecast runs:

EX 2.2 Retrieving initialized DWD analysis data:

- Request native analysis data from DWD’s meteorological data management system SKY for the date 2017-06-01T00:00:00 at a horizontal grid spacing of 13 km using the PAMORE tool, see Section 2.2.1:

  ```
  pamore -d date -hst 0 -htp 0 -lt a -model iglo -iglo_startdata_0
  ```

- Take a look at the data with the grib ls command-line tool.
C. Exercises

C.2 Necessary Input Data

- Export the environment variable `GRIB_DEFINITION_PATH` with the setting
  
  \[
  \text{GRIB\_DEFINITION\_PATH} = \text{/usr/local/pkg/grib\_api/definitions/} \ldots \text{release/$\text{GRIB\_API\_VERSION}/definitions}
  \]

  and run `grib ls` once more. Are there any differences w.r.t. the displayed short names? See Section 1.1.2 for an explanation.

Remapping of initial data:

- Create a copy of the run script
  
  `dwd_icon_tools/icontools/xce_remap_inidata`

  which performs the task of remapping the variables of an uninitialized analysis product (Table 2.2).

- Adapt the set of variables which is to be remapped for initialized analysis products, according to Table 2.3, i.e. replace the fields `VN`, `THETA_V`, `DEN` by the fields `T`, `U`, `V`, `P`.

- Submit the script to the batch system. Afterwards repeat the remapping procedure for the 13 km nest (R3B07) over Europe.

- Copy and rename the remapped analysis files s.t. the file names match the nomenclature `dwd\_ANA_R<nroot>B<jlev>_DOM<idom>.grb`, see Section 5.1.2.

In this exercise we will prepare the initial data for ICON to start from an IFS analysis:

Retrieving IFS data:

- **Download** – *This step has to be executed on the ECMWF site:*
  Start the `mars4icon` script and download an IFS analysis file for the date 2017-01-12T00:00:00, see Section 2.2.2.

- Interpolate the data horizontally from the lat/lon grid onto the triangular ICON grid using `iconremap`. The run script
  
  `dwd_icon_tools/example/runscripts/xce_ifs2icon.run`

  will do this job.

  Fill in the name of the IFS file by setting `in\_grid\_filename` and `in\_filename`. Further help on `iconremap` can be found in Section 2.2.3.

- Submit the `iconremap` job to the Cray XC 40.

- List the fields contained in the output file using `cdo` and/or `ncdump`. Compare this to Table 2.4. Besides, find out which field(s) are not defined on cell circumcenters.
## Preparation of Limited Area Runs

**Note:**

The following exercise requires a number of data files which are produced as model output in the real data exercise, Ex. C.5.5.

In this exercise we will pre-process the initial and boundary data for ICON limited area runs:

### Local grid file and its external parameters:

- In the directory `case_lam/input` you find an archive file which is the result of the web-based grid generator described in Section 2.1.4.

  Uncompress this file and visualize its content with the NCL file `case_lam/plot_grid.ncl`.

  See Figure C.3 for a reference (without highlighted boundary).

- Investigate the grid file with the `ncdump` utility (see Section 9.1.1): How many triangle cells are contained in the local grid? – **Answer:** __________ cells

### Remapping of initial data:

The directory “`lam_forcing`” generated by the real case run contains a file with the prefix `init`, which will serve as initial conditions for the limited area run. Remap the data onto the local grid.

The subdirectory `case_lam` contains a script `xce_remap_inidata` which will do the remapping.

- Insert the path to the ICON Tools binaries, and
- adapt the path to the input data file (directory “`lam_forcing`” generated by real case run).

Inspect the local (TARGET) grid file and the grid which was used in Ex. C.5.5 for creating the initial data (SOURCE).

- Identify the grid spacing of both grids in ICON’sRxBy nomenclature. (You could make use of `ncdump`.)

<table>
<thead>
<tr>
<th>SOURCE grid</th>
<th>remap data</th>
<th>TARGET grid</th>
</tr>
</thead>
<tbody>
<tr>
<td>__________</td>
<td>__________</td>
<td>__________</td>
</tr>
</tbody>
</table>

- If the source grid has a horizontal grid spacing of \( \approx 13 \text{ km} \), what is the grid spacing of your local (TARGET) grid in km, given the RxBy expressions identified above (see Eq. 2.1)?

  – **Answer:** TARGET grid res. __________ km

Submit the script to the batch system.

- Check the result: The remapping script should have created a file with prefix `init` in `case_lam/output`. 
C.4 Running Idealized Test Cases

In this exercise you will learn how to set up idealized runs in ICON with and without nested domains.

Job submission to the Cray XC 40 can be performed on the Linux cluster lce. Note, however, that visualization tools (CDO, NCL, ncview) are only available on the lce!

Running the Jablonowski-Williamson Test Case

In this exercise we will set up the Jablonowski-Williamson baroclinic wave test in ICON:

Preparations:

Retrieve the necessary grid file from the ICON download server (see Section 2.1.2):

- Open the download page for the pre-defined ICON grids http://icon-downloads.mpimet.mpg.de in your web browser.
- Pick the R2B05 grid no. 14 from the list and right-click on the hyperlink for the grid file, then choose "copy link location".

Remapping of boundary data:

- The subdirectory case_lam contains a copy of the DWD ICON Tools script xce_limarea, see Section 2.3. Open this script and
  - insert the path to the ICON Tools binaries,
  - adapt the path to input data files (directory “lam_forcing” generated by real case run).
- Submit the script to the batch system.
- Check the result: Visualize the boundary data with the NCL script case_lam/plot_boundary_data.ncl.
- Investigate the files: How many cells are contained in the boundary grid?
  - Answer: [ ] cells
- When looking at the set of boundary data variables, do you have any idea for further improvement in terms of storage space?
  - Answer: [ ]
C. Exercises

Open a terminal window, login into the Linux cluster lce, and change into the subdirectory `test_cases/case_idealized/input`. Download the grid file into this subdirectory by typing:

```
wget link_location -e http-proxy=ofsquid.dwd.de:8080.
```

Run the Jablonowski-Williamson (JW) test case without nested domains:

- Change into the run script directory `test_cases/case_idealized`. The run script is named `run_ICON_R02B05_JW`.
- Fill in the name of the ICON model binary (including the path) and several missing namelist parameters. I.e. set `ltestcase`, `ldynamics`, `iforcing`, `nh_test_name` and `itopo`. See Section 1.1.1 for the location of your ICON model binary and Section 4.2.1 for additional help on the namelist parameters.
- Submit the job to the Cray XC 40, using the PBS command `qsub`.
- Check the job status via `qstat` and `qcat` (see Section A).

Inspecting the output:

- Go to the output directory of `case_idealized`. You will find three NetCDF output files with (a) model level output on the native (triangular) grid, (b) pressure level output on the native grid, (c) model level output interpolated onto a regular lat-lon grid.
- Have a closer look to the different output files and their internal structure to find out which one is which. We suggest to use the tools `cdo` and `ncdump` as described in Section 9.1.
- What output interval (in h) has been used for model level output on the triangular grid?  
  - **Answer:** [ ] h

- Visualize the output with one of the tools described in Section 9. An NCL script named `JW_plot.ncl` is available in `test_cases/case_idealized`.
- Compare the output of the NCL script with the reference `JABW_DOM01.ps` given in the subdirectory `reference`.
- Modify the NCL script by setting `lfocus_sh=True` and re-run. This will basically re-create the first set of plots, but with a focus on the southern hemisphere and different contours.
- Is the flow still purely zonal after 9 days? Describe what you see. Do you have an explanation for the observed behavior?  
  - **Answer:** [ ]
C.4 Running Idealized Test Cases

Nested Sub-Domains

In this exercise we will repeat the run of the previous exercise (Jablonowski-Williamson baroclinic wave test) with a nested sub-domain:

- Go to the run script directory. The run script for a nested grid experiment is termed `run_ICON_R02B05N6_JW`.
- Fill in the name of your ICON model binary and missing namelist parameters. I.e. extend
  - `dynamics_grid_filename`,
  - `num_lev (run_nml)` and
  - `dynamics_parent_grid_id (grid_nml)`, see Section 4.2.2 for additional help. The same output fields as for the global domain will be generated for the regional domain(s).

The grid files of the nested domains have already been prepared for you. They are stored in the subdirectory `test_cases/case_idealized/input`. The location of the prepared nests is depicted in Figure 4.3. Feel free to add one or both nests to your global domain.

- Submit the job to the Cray XC 40.
- After the job has finished, visualize the output on the global domain as well as on nest(s) using one of the tools described in Section 9. When using NCL (`JW_plot.ncl`), you will have to adapt `workdir` and `domain_nr`.
- Compare the results on the global domain with those of the previous exercise. Does the global domain output differ? If so, do you have an explanation for this?
  - Answer: 

Additional question: setting the parent grid ID.

The figure to the right depicts an alternative nest configuration, in which two domains are nested into each other. How does the correct setting for `dynamics_parent_grid_id` look like, given that the grid files are ordered as follows:

```
dynamics_grid_filename = 'gridFileA','gridFileB','gridFileC'
```

- Answer: `dynamics_parent_grid_id = `
Optional Exercise: Tracer Advection

The JW test case provides a set of four pre-defined idealized tracer fields (see Figure 4.2). Here, you will learn how to enable and control the transport of passive tracers in idealized tests. See Section 3.6.5 for details on the configuration of the tracer transport for standard NWP runs.

- Enable tracer advection:
  - Go to the directory test_cases/case_idealized and open the run script run_ICON_R02B05N6_JW.
  - Enable the transport module by setting the main switch ltransport to .TRUE..
  - Enable (uncomment) the namelist transport_nml, which specifies details of the applied transport scheme.
  - Select one or more tracers from the set of pre-defined tracer distributions (see Figure 4.2). A specific distribution can be selected by adding the respective tracer number (1, 2, 3, or 4) to tracer_inidist_list (namelist nh_testcase_nml, comma-separated list of integer values).

- Extend the output namelist
  - Add the selected tracer fields to the list of output fields in the namelists output_nml (namelist parameters ml_varlist and pl_varlist).
  
For idealized test cases, tracer fields are named q_x, where x is equal to the suffix specified via the namelist variable tracer_names (namelist transport_nml, comma-separated list of string parameters). The n^{th} entry in tracer_names specifies the suffix for the n^{th} entry in tracer_inidist_list. If nothing is specified, tracer fields are named q_x, where x is a number indicating the position of the tracer within the ICON-internal 4D tracer container.

- Visualization: Enable lplot_transport in your NCL script. You can also have a quick look using ncview.

- Have a look at tracer q_4. Initially it is constant (= 1) everywhere. Ideally, an initially constant tracer should stay constant for all times, no matter how complicated the flow.
  Does ICON preserve a constant tracer in a run without nest?

  - Answer: 

  Does ICON preserve a constant tracer in a run with nest?

  - Answer: 

- Avoid non-physical negative values for tracer q_1.
C. Exercises

C.5. Running Real Data Test Cases

In this exercise you will learn how to start ICON from DWD analysis data and how to perform a multi-day forecast with 26 km grid spacing globally and 13 km over Europe. Another practical aim of this exercise is to create raw data for driving a limited area ICON run. Further use of this data will be made in Ex. C.6.1.

The configuration and compilation of the ICON code has to be done on the Cray XC 40 login node `xce`. Job submission to the Cray XC 40 can be performed on either the login node `xce` or the Linux cluster `lce`. **Note, however, that visualization tools (CDO, NCL, ncview) are only available on the Linux cluster `lce`!**

**Input Data**

The exercises in this section require a number of grid files, external parameters and input data. This data is already available in the directory `case_realdata/input`. Their creation process is explained in Ex. C.2.1 and C.2.2 (see p. 218).

On default, ICON expects the input data to be located in the experiment directory (termed `$EXPDIR` in the run scripts). The run script creates symbolic links in the experiment directory, which point to the input files.

**Note:**

Note that the names originating from the database requests (see Section 2.2.1) differ from the file names above. The input files have to be renamed in this case in order to match the default filename structure expected by the model.
Starting a Global ICON Forecast from DWD Analysis

In this exercise you will learn how to run ICON in real data mode:

Open the ICON run script `case_realdata/run_ICON_R03B06_dwdini` and prepare the script for running a global **72 hour forecast** on an R3B06 grid with 26 km horizontal grid spacing without nest. The start date is

2017-01-12T00:00:00 , i.e. January 12, 2017.

**Basic settings**

- Fill in the missing namelist parameters `ini_datetime_string`, `end_datetime_string`, `ltestcase`, `ldynamics`, `ltransport`, `iforcing`, and `itopo` for real data runs (see Section 5.1.1).

- For better runtime performance, switch on asynchronous output by setting the number of dedicated I/O processors (`num_io_procs`) to a value larger than 0 (e.g. 1). See Section 7.3.1 for more details on the asynchronous output module.

**Specifying the input data**

- The grid file(s) to be used are already specified in the run script (see `dynamics_grid_filename`, `radiation_grid_filename`).

- Note that – funnily enough – the initialized analysis file is provided via the namelist parameter `dwdfg_filename`, see Section 5.1.4.

Due to the appropriate choice of the file names in the experiment directory, ICON is able to locate the analysis data sets automatically without specifying the namelist variable `dwdfg_filename`. However, it is given in the run script for reference, using the keyword nomenclature mentioned in Section 5.1.2.

Have a look at these settings and try to understand how these keywords work.

Which of the following filenames would be accepted by the ICON model?

- `dwdANA_R3B06.grb`
- `dwdANA_R2B6_DOM01.grb`
- `dwdANA_R9B02_DOM02.grb`
- `dwdana_R2B06_DOM01.grb`

- Specify the name of the external parameter file (`extpar_filename`). Instead of specifying the full name, try to make use of the keyword `idom`.

**Settings for the Initialized Analysis Data**

- Choose the appropriate initialization mode `init_mode` for starting the model from DWD initial analysis data (see Section 5.1.4).
C.5 Running Real Data Test Cases

- Surface tiles are activated in the given run-script, as can be deduced from the parameter `ntiles` in the namelist `ind_nml`. Since the initial data contain aggregated (cell averaged) surface fields, a tile coldstart must be performed by setting `ltile_coldstart` appropriately, i.e. each tile must be initialized with the same cell averaged value (see Section 3.8.10).

Running the model and inspecting the output

- Submit the job to the Cray XC 40.
- After the job has finished, inspect the model output:
  - Take a look at the output files in `case_realdata/output`. You should find two files named `NWP...`. Use `cdo sinfov` to identify the
    - time interval between two outputs – Answer: ______ h
    - total time interval for which output is written – Answer: ______ h
    - type of vertical output grid (ML, PL, HL) – Answer: ______
  - one file contains output on the native ICON grid, the other one output on a regular lat/lon grid. Use `cdo sinfov` to identify which file is which.
    - Answer: native
    - Answer: lat/lon
  - Visualize the 2 m temperature, integrated water vapour, gusts at 10 m, and total precipitation using the `ncview` utility. If you like, you can look into other fields as well.

Time-stepping

This exercise focuses on aspects of the ICON time-stepping scheme, explained in Section 3.7.1:

- Compute the `dynamics` time step $\Delta \tau$ from the specification of the physics time step $\Delta t$ (dtime) and the number of dynamics substeps `ndyn_substeps`.
  
  - Answer: $\Delta \tau = ______$ s

- Take a look at Equation (3.39) and calculate an estimate for the maximum dynamics time step which is allowed for the horizontal grid spacing at hand.
  
  - Answer: $\Delta \tau_{\text{max}} = ______$ s

Now compare this to the time step used: Did we make a reasonable choice?
Parallelization and Run-Time Performance

In this exercise you will learn how to specify the details of the parallel execution. We will use the ICON timer module for basic performance measuring.

**EX 5.3**

Performance assessment using the timer output:

- Open your run script from the previous Exercise C.5.1 and enable the ICON routines for performance logging (timers). To do so, follow the instructions in Section 7.3.4.
- Repeat the model run.
- At the end of the model run, a log file is created. It can be found in your base directory `case_realdata`. Scroll to the end of this file. You should find wall clock timer output comparable to that listed in Section 7.3.4. Try to identify the total run-time – Answer: __________ s
  
  the time needed by the radiation module – Answer: __________ s

This exercise focuses on the mechanisms for parallel execution of the ICON model. These settings become important for performance scalability when increasing the model resolution and core counts:

**EX 5.4**

Changing the number of MPI tasks: In case your computational resources are sufficient, one possibility to speed up your model run is to increase the number of MPI tasks.

- Create a copy of your run script `run_ICON_R03B06_dwdini` named `run_ICON_R03B06_dwdini_fast`. In order to avoid overwriting your old results, replace the output directory name (EXPDIR) by `exp02_dwdini_fast`.
- Double the total number of MPI tasks compared to your previous job and re-submit. In more detail, do the following:
  - Your old script (run_ICON_R03B06_dwdini) ran the executable in hybrid mode on 25 nodes using 12 MPI tasks/node and 4 OpenMP threads/MPI task with hyper-threading enabled.
  - Your new script (run_ICON_R03B06_dwdini_fast) should run the executable in hybrid mode on 50 nodes using 12 MPI tasks/node and 4 OpenMP threads/MPI task with hyper-threading enabled.

  You need to adjust both the PBS settings and the `aprun` command. See Section 7.3.4 for additional help.
- Compute the speedup that you gained from doubling the number of MPI tasks.
C.5 Running Real Data Test Cases

Figure C.1.: Computational grid with a two-way nested region over Europe (yellow shading). The outline of the COSMO-EU domain (formerly used operationally by DWD) is shown in red for comparison.

- Compare the timer output of the dynamical core, \texttt{nh\_solve}, and the transport module, \texttt{transport}, with the timer output of your previous run.

\begin{tabular}{|c|c|c|c|}
\hline
\texttt{nh\_solve} & \texttt{nh\_solve} & \texttt{transport} & \texttt{transport} \\
\texttt{25 nodes} & \texttt{50 nodes} & \texttt{25 nodes} & \texttt{50 nodes} \\
\hline
\hline
s & s & s & s \\
\hline
\end{tabular}

What do you think is a more sensible measure of the effective cost: total \texttt{min} or total \texttt{max}?

- \textit{Answer: }

Which speedup did you achieve and what would you expect from “theory”?

- \textit{Answer: } Speedup achieved = \frac{T_{\text{50 nodes}}}{T_{\text{25 nodes}}} =

- \textit{Answer: } Speedup expected =

\[ \text{Speedup achieved} = \frac{T_{\text{50 nodes}}}{T_{\text{25 nodes}}} \]
Writing Output for Driving a Limited Area Simulation

In this exercise you will learn about some of the output capabilities of ICON. We will set up a new output namelist and generate a data set which enables us to drive a limited area version of ICON as described in Chapter 6.

In order to achieve a somewhat higher spatial resolution at acceptable costs, we will switch on a two-way nested region over Europe with a horizontal grid spacing of 13 km (see Figure C.1):

- Create a copy of your run script runICON_R03B06_dwdini named runICON_R03B06N7_dwdini. Alternatively, you may adopt the settings of the reference version reference/case2/runICON_R03B06_dwdini.
- In order to avoid overwriting your old results, replace the output directory name (EXPDIR) by exp02_dwdini_lamforcing.
- Activate the nest, by extending the list of horizontal grids to be used (dynamics_grid_filename) and the parent grid IDs (dynamics_parent_grid_id (grid_nml)).
- In order to save some computational resources, the nested region should have a reduced model top height and comprise only the lowermost 60 vertical levels of the global domain (instead of 90 levels). Please extend num_lev (run_nml) accordingly.

Adding a new output namelist:

- The run script contains two commented-out output namelists. One is meant for writing forcing (boundary) data, the other for writing initial data for driving a limited area run. Activate the namelists and fill in the missing parameters. See Section 7.1 for additional details regarding output namelists. Forcing data should be written
  - in GRIB2 format
  - for the EU-nest only
  - 2-hourly from the start until 48 hours forecast time
  - with one output step per file
  - on the native (triangular) grid
  - into the subdirectory “lam_forcing” of your output directory exp02_dwdini_lamforcing, using the filename prefix “forcing”. If the subdirectory does not exist, please create.
  - containing model level output for U, V, W, PRES, TEMP, QV, QC, QI, QR, QS, HHL.
C.5 Running Real Data Test Cases

– Take a look at the available ICON variable groups listed in Appendix B. Is there an output group that can be used instead of the list of variables? If yes, which group?
  – Answer:

– Is there also an ICON variable group that writes all variables needed to initialize an ICON-LAM simulation? If yes, which group?
  – Answer:

• Submit the job to the Cray XC 40.

Check the correctness of your output files:

• You should find 25 files in your output directory “lam_forcing”. Apply the command `cdo sinfov data-file.grb > data-file.sinfov` to the last file. Compare with the reference output in Table C.1 to see whether your output namelist is correct.

Optional: A Deeper Look into Spin-Up Effects

Depending on the data set used to initialize the model, spin up effects may become visible during the first few hours of a forecast. This is especially true if third party (i.e. non-native) analysis data sets are used. Here we will have a look into the spin up behavior when ICON is started from native vs. non-native analysis. As an example for a popular non-native analysis, we will choose data from the IFS.

• Run the NCL script `case_realdata/water_budget.ncl`, to get a deeper insight into the model’s spin up properties and water budget when started from DWD analysis. The script generates time series of vertically integrated water vapour $tqv$ and condensate $tqx$ from the model level output in `case_realdata/output/exp02_dwdini`.

  Compare the results to Figure C.2 which shows the corresponding results when ICON is started from IFS analysis. Note that Figure C.2 has been produced for a longer time range.

  Which analysis data set leads to an increased spin up/down in this particular case?
  
  native analysis (DWD)  
  non-native analysis (IFS)

  The additional NCL plots will give you some insight into the global atmospheric water budget

  \[
  \frac{dQ_t}{dt} = P - E + R ,
  \]
Table C.1.: Reference output for Ex. C.5.5. Structure and content of file forcing_ML_20170114T000000Z.grb

<table>
<thead>
<tr>
<th>File format : GRIB2</th>
</tr>
</thead>
<tbody>
<tr>
<td>-1 : Institut Source Type</td>
</tr>
<tr>
<td>1 : DWD unknown instant</td>
</tr>
<tr>
<td>2 : DWD unknown instant</td>
</tr>
<tr>
<td>3 : DWD unknown instant</td>
</tr>
<tr>
<td>4 : DWD unknown instant</td>
</tr>
<tr>
<td>5 : DWD unknown instant</td>
</tr>
<tr>
<td>6 : DWD unknown instant</td>
</tr>
<tr>
<td>7 : DWD unknown instant</td>
</tr>
<tr>
<td>8 : DWD unknown instant</td>
</tr>
<tr>
<td>9 : DWD unknown instant</td>
</tr>
<tr>
<td>10 : DWD unknown instant</td>
</tr>
<tr>
<td>11 : DWD unknown instant</td>
</tr>
</tbody>
</table>

Grid coordinates:
- unstructured: points=172740
- uuid: 890df70c-a566-3b91-438f-f527deb82760

Vertical coordinates:
1: generalized_height: levels=60
   - height: 1.5 to 60.5 by 1
   - bounds: 1-2 to 60-61 by 1
   - zaxis: number = 4
   - uuid: a0919ba7-df84-ce2b-c4c2-3c215c033520

2: generalized_height: levels=61
   - height: 1 to 61 by 1
   - zaxis: number = 4
   - uuid: a0919ba7-df84-ce2b-c4c2-3c215c033520

3: generalized_height: levels=61
   - height: 0.5 to 30.5 by 0.5
   - bounds: 1-0 to 61-0 by 0.5
   - zaxis: number = 4
   - uuid: a0919ba7-df84-ce2b-c4c2-3c215c033520

Time coordinate: 1 step
- RefTime = 2017-01-12 00:00:00
- Units = minutes
- Calendar = proleptic_gregorian
- YYYY-MM-DD hh:mm:ss YYYY-MM-DD hh:mm:ss YYYY-MM-DD hh:mm:ss YYYY-MM-DD hh:mm:ss
- 2017-01-14 00:00:00

where $Q_t$ is the vertically integrated atmospheric water content and $\frac{dQ_t}{dt}$ is the rate of change over time. $P$ is the amount of total precipitation, $E$ is the total evaporation, and $R$ is a residuum. Is the budget closed (i.e. is $R = 0$ in your model run)?
C.5 Running Real Data Test Cases

Figure C.2.: Time series of area averaged column integrated specific moisture $\langle tqv \rangle$ (top) and condensate classes $\langle tqx \rangle$ (bottom) for a 7-day forecast started from IFS analysis fields. Start date was 2017-01-12T00:00:00. A spin up in $\langle tqv \rangle$ and initial adjustments in $\langle tqx \rangle$ is clearly visible.
Optional: Checking for Bit-Reproducibility

In this exercise we test the ICON model for bit-reproducibility, i.e. the feature that running the same binary multiple times results in bitwise identical results.

- Compare the model level output of `run_ICON_R03B06_dwdini` with the output that was produced by your modified script `run_ICON_R03B06_dwdini_fast`. You can use `cdo infov data-file.nc` for getting information about the contents of your output file. You should dump this information into text files:

  ```
  cdo infov data-file.nc > data-file.infov
  ```

  so that you can compare them later on. Due to the limited number of digits printed by CDO, this is no check for bit-reproducibility in a strict mathematical sense, however experience showed that `cdo infov` is very reliable in revealing reproducibility issues.

- Now compare the model level output produced by your modified script `run_ICON_R03B06N7_dwdini` with the output produced by `run_ICON_R03B06_dwdini`. Is the result different from the previous comparison? Do you have an explanation for this?

  **Hint:** For a convenient comparison of ASCII files you may use the `tkdiff` utility.

C.6. Running ICON-LAM

The exercises in this section require a number of grid files, external parameters and input data. In particular, initial data and driving boundary data are required. Both were produced in the real data exercise, Ex. C.5.5. The preprocessing of this data is explained in Ex. C.2.4 (see p. 220).

Running ICON in Limited Area Mode

In this exercise we will run ICON in limited area mode. The model will be driven by initial and boundary data which have been produced in Ex. C.2.4.

Open the run script `case_lam/run_ICON_R3B08_lam` and prepare it for running a **48 hour forecast** on a **limited area grid over Germany** (see Figure C.3). As for the global run, the start date is

```
2017-01-12T00:00:00 , i.e. January 12, 2017.
```

The run script is geared up insofar as all soft-links which specify the model binary, grids, initial and boundary data are already set. Your main task will be to set up the ICON namelists for a limited area run.
C.6 Running ICON-LAM

Figure C.3.: Illustration of the local grid used in Exercises C.6.1–C.8.3. The horizontal grid spacing is ≈ 6.5 km (which corresponds to R3B08 in ICON nomenclature). The boundary region is highlighted, where nudging towards the driving data is performed. The driving data for this test case has been created in Ex. C.2.4.

Basic settings:
- Set the correct start and end date:
  
  `ini_datetime_string`, `end_datetime_string`.

- Switch on the limited area mode by setting `l_limited_area` and `init_mode` (see Section 6.3).

Initial data:
- Specify the initial data file via `dwdfg_filename`. Since we do not make use of additional analysis information from a second data file, remember to set `lread_ana` accordingly (see Section 6.3).

- Specify a dictionary file for the initial data via `ana_varnames_map_file` (see Section 5.1.1).
  
  **Hint:** Take a closer look at `case_lam/run ICON_R3B08_lam`, where a link to the correct dictionary is already set.
C. Exercises

The dictionary has the purpose to translate the variable names in the GRIB2 or NetCDF input file into the ICON internal variable names. According to the dictionary used for this exercise, what are the internal variable names and the GRIB2/NetCDF variable names for

- **Pressure:**
  - GRIB2/NetCDF: [Blank]
  - ICON internal: [Blank]

- **Temperature:**
  - GRIB2/NetCDF: [Blank]
  - ICON internal: [Blank]

- **Half level height:**
  - GRIB2/NetCDF: [Blank]
  - ICON internal: [Blank]

**Boundary data:**

- Specify the lateral boundary grid and data via `latbc_boundary_grid`, `latbc_path` and `latbc_filename`. For the latter you will have to make use of the keywords `<y>`, `<m> <d>` and `<h>` (see Section 6.4.1).

- What is the time interval between two consecutive boundary data files? The command `cdo sinfov` may be helpful.
  - **Answer:** s
  - Set the namelist parameter `dtime_latbc` accordingly.

- Inspect the number of vertical levels in a boundary data file. Is it different from the number of vertical levels that is used by the model itself?
  - **Answer:** [Blank]

**Running the model and inspecting the output**

- Extend the lat-lon output namelist by the 2m temperature, surface pressure, mean sea level pressure, and 10 m gusts. See Appendix B for variable names and description.

- Submit the job to the Cray XC 40.

- After the job has finished, inspect the model output. You should find multiple files in the output directory `case.lam/output/exp03_R3B08_dwdlam` which contain hourly output on model levels remapped to a regular lat-lon grid.
  - Take a look at the output fields by using `ncview`. How would you characterize the overall weather situation for that time period?
    - southfoehn over the alpine region [Blank]
    - strong frontal system passing over Germany [Blank]
    - weak frontal system passing over Germany [Blank]
    - anticyclonic situation with very calm winds [Blank]
Temporal Resolution of the Boundary Data

By playing around with the temporal resolution of the boundary data you will get some idea how this might affect your simulation results.

Create a copy of your run script `run_ICON_R3B08_lam` and name it `run_ICON_R3B08_lam_lowres`. Replace the output directory name (EXPDIR) by `exp03_R3B08_dwdlam_lowres` in order to avoid overwriting your results.

- Halve the time resolution of your forcing (boundary) data, see Section 6.3 for the namelist parameter. Write down your chosen value:
  
  - *Answer:* [ ] s

- Submit the job to the Cray XC 40.

- Compare the results with your previous run. Does the time frequency with which the boundary data are updated have a significant impact on the results? You can visualize cross sections with the NCL script `case_lam/plot_cross_section.ncl` and/or make use of ncview.

Driving ICON-LAM by DWD Data

This exercise is similar to the previous exercise C.6.1 inasmuch as we will run ICON in limited area mode again. However, rather than using initial and boundary data that have been created in one of the previous exercises, we will initialize and drive ICON-LAM by official DWD analysis and forecast data. This setting might be closer to what you will encounter when you run ICON-LAM at your home institution.

The data products for this particular exercise will be provided. Note, however, that PAMORE data requests in general require registration via datenservice@dwd.de.

In the subdirectory

```
/e/uwork/trng024/packages/pamore_data
```

you will find the raw data for an ICON limited area run consisting of an initialized analysis and two-hourly forecasts. This data set has been downloaded from the DWD database with the help of the PAMORE tool, see Section 2.2.1 for details.

- For the sake of simplicity, link the above directory into `case_lam/input_dwd`.
- Take a look at the data with the `grib_ls` and/or `cdo` command-line tool. Find out about the analysis date and the time range of the data set.
- *Imagine that your boss at home asks you to retrieve this data set from the DWD database.*
Write down the PAMORE commands which you would have to insert into the PAMORE web form. See Section 2.2.1 and 2.3 for help.

- **Analysis:**

- **Forecasts:**

  - Export the environment variable `GRIB_DEFINITION_PATH` with the setting

    ```
    GRIB_DEFINITION_PATH=/usr/local/pkg/grib_api/definitions/... release/$GRIB_API_VERSION/definitions
    ```

  and run `grib ls` once more. Are there any differences w.r.t. the displayed short names? See Section 1.1.2 for an explanation.

**Remapping of initial data:**

In order to use the DWD initialized analysis for your LAM run, it must be remapped to your limited area grid.

- Determine the source grid on which the raw data are defined. Write down the grid identifier `numberOfGridUsed`.
  
  - **Answer:** `numberOfGridUsed` =

- Retrieve the matching grid file from the ICON download server (see Section 2.1.2):
  
  - Open the download page for the pre-defined ICON grids [http://icon-downloads.mpimet.mpg.de](http://icon-downloads.mpimet.mpg.de) in your web browser.
  
  - Pick the matching grid number from the list and right-click on the hyperlink for the grid file, then choose ”copy link location”.
  
  - Open a terminal window, login into the Linux cluster `lce`, and change into the subdirectory `case_lam/input_dwd`. Download the grid file into this subdirectory by typing

    ```
    wget link_location -e http-proxy=ofsquid.dwd.de:8080
    ```

  - Write down the resolution in ICON’s RxBy nomenclature.
    
    - **Answer:** `RxBy` =

- Create a copy of the remapping script `xce_remap_inidata` located in the subdirectory `case_lam`.

- Open the script and adapt
  
  - the path to the input data file that has been retrieved from the DWD database (directory `case_lam/input_dwd/pamore_data`).
  
  - the file name of the input grid.
  
  - the output directory such as not to overwrite your previous initial data. You might store your output in a new sub-directory `output_dwd` located inside the base directory ”`case_lam`”.
C.6 Running ICON-LAM

- Submit the script to the batch system.

**Remapping of the lateral boundary conditions:**

In contrast to the previous ICON-LAM exercises we will interpolate the forcing data onto the full limited area grid instead of the lateral boundary strip only. Can you think of simulations where such a setting is required?

- **Answer:**

Create a copy of the remapping script for boundary data *xce_limarea*:

- Open this script and adapt the path to your input data files (directory “case_lam/input_dwd/pamore_data”), the file name of the input grid as well as the output directory (“case_lam/output_dwd”).

- Adapt the script such that the AUXGRID auxiliary grid (boundary strip) is no longer used but the full limited area grid (LOCALGRID).

Submit the script to the batch system.

**Small consistency check:** How many grid cells do you expect in the output files?

- **Answer:** _cells_

**Running the model**

Create a copy of the run script *case_lam/run_ICON_R3B08_lam* and prepare it for running a 36 hour forecast on the same limited area grid over Germany as used in Ex. C.6.1 (see Figure C.3). In order to avoid overwriting your old results, replace the name of the output directory.

**Initial and boundary data:**

As we will use a new set of initial and boundary data, several changes are necessary:

- Change the settings for the analysis input files (initial data).

- Specify the lateral boundary data via `latbc_path` and `latbc_filename`. For the latter you will have to make use of the keyword `<ddhhmmss>` (see Section 6.4.1).

- Check whether the time interval between consecutive boundary data files is set correctly (`dtime.latbc`).

Submit the job to the Cray XC 40.

Did your run finish successfully?
C. Exercises

C.8. Programming ICON

Allocating Additional Fields

In this exercise we learn how to allocate new ICON variables. As a by-product we will visualize ICON’s domain decomposition.

This requires to modify the ICON code, where details are given in Section 8.3. Remember to create a fully copy of your src directory. Subsequent exercises will be based upon the unmodified sources!

Step-by-step checklist

In the subdirectory src/atm_dyn_iconam:

a) Open the module mo_nonhydro_types.
   Insert a new 2D variable pointer process_id of type REAL(wp) to the derived type TYPE(t_nh_diag).

b) Open the module mo_nonhydro_state.
   At the end of the subroutine new_nh_state_diag_list:
   initialize meta-data variables for NetCDF and also for GRIB2 with discipline = parameterCategory = parameterNumber = 255
   and place the add_var call for the new field.

c) Now, after the add_var call, the new field has been allocated.
   Fill the data array with a constant value:
   ```
   process_id(:,:, :) = get_my_mpi_work_id()
   ```
   This auxiliary function from mo_mpi returns the MPI rank of the processor (see Section 8.2.4).

d) Open the namelist of the previous test case C.6.1 and insert the new fields in the lat-lon output specification.

Compile and run the model.
Visualize the results (ncview).

Please answer the following questions:

- Why is the output field not a constant value?
  - Answer: 

- Take a closer look at the data: Can you guess why the output field does not only contain integer values?
  - Answer: 
In this exercise we will implement the technical setup for the calculation of new diagnostic fields:

In this exercise we will implement the technical setup for the calculation of the following new diagnostic fields (2D variables):

- **RHI\_MAX**: Relative humidity over ice (hourly maximum in the column, [%])
- **QI\_MAX**: Maximum cloud ice content (hourly max. in the column, $[\text{kg/m}^2]$)

Again, this requires to modify the ICON code.

- Repeat the steps a) – b) of Exercise C.8.1 and allocate two new 2D variables RHI\_MAX, QI\_MAX.
- Open the module mo\_nh\_stepping (subdirectory src/atm\_dyn\_iconam).
  - Create an (empty) subroutine calculate\_diagnostics and place a call to this subroutine immediately before the call to the output routine.
- Fill your subroutine with a 2D loop over all prognostic grid points, see p. 161 for help.
  - Use this loop to initialize the fields to a constant value.
    
    \[
    p\_nh\_state(jg)\%diag\%RHI\_MAX(jc,jb) = 2.0\_wp \\
    p\_nh\_state(jg)\%diag\%QI\_MAX(jc,jb) = 2.0\_wp
    \]
    
  - All other values should be set to 1.0\_wp.

How many triangle rows do you expect to have the value 1 in the output?

- Answer: ______ rows.

- Open the namelist of the previous test case C.6.1 and insert the new fields in the lat-lon output specification.
- Compile and run the model. Visualize the results (ncview).
- **Final question**: Instead of the value 2.0\_wp we’d now like to initialize RHI\_MAX with the lower-most level of the (diagnostic) temperature field. How could we achieve this?

- Answer: ______
Implementing New Diagnostics

In this exercise we will calculate the new 2D diagnostic fields $$\text{RHI}_{\text{MAX}}$$, $$\text{QI}_{\text{MAX}}$$:

- Modify the subroutine `calculate_diagnostics`:
  Instead of initializing the fields to a constant value as in Ex. C.8.2, add a loop over the vertical model levels for every prognostic grid point.

- Then fill your subroutine with the calculation of the two new quantities. Three hints:
  - The module `mo_util_phys` (subdirectory `src/atm_phy_nwp`) may offer some help with respect to $$\text{RHI}_{\text{MAX}}$$.
  - Exner pressure field: values for domain “jg” are accessed via `p_nh_state%prog(nnow(jg))%exner(:, :, :)`.
  - 3D tracer field QI (same for QV): values for domain “jg” are accessed via `p_nh_state%prog(nnow_rcf(jg))%tracer_ptr(iqi)%p_3d(:, :, :)`. The constants iqv, iqi can be found in `mo_run_config`.

- Revisit the `add_var` calls of the quantities $$\text{RHI}_{\text{MAX}}$$, $$\text{QI}_{\text{MAX}}$$: specify an hourly reset, see p. 162 for an explanation.

- Open the namelist of the previous test case C.6.1 and insert the new fields in the output specification.

Compile and run the model. Visualize the results (`ncview`).
C.10. Running ICON-ART

In order to start with the exercises, you have to copy and unpack the ART code inside your ICON source directory as described in section 10.3.

You will find the ART code at: 
/e/uwork/trng024/packages/art.tar.gz.
Folders with the input data for all ART exercises can be found at: 
/e/uwork/trng024/packages/ART-INPUT.

After you have copied the source code, you have to install ICON-ART. For this purpose, proceed as described in Section 10.3. After a successful compilation of ICON-ART, you can start with the experiments, that were prepared for this purpose:

Point Sources in ICON-ART LAM

In this exercise, you will learn to add your own tracers with emissions from point sources. You are free to choose the tracer(s) to transport and the location of the point source(s).

In order to perform the simulation, you have to do the following steps:

• Inside $ARTDIR/runctrl_examples/run_scripts you will find the run script for this test case called exp.art.trng19.case1.pntSrc
• inside the ART-INPUT folder you will find a folder called CASE1-PNTSRC containing all input data required for this test case
• Edit the run script according to the namelist parameters you find in Section 10.4. You will find a ? at all places where you have to edit something.
• Create the XML files that are needed to define tracers (see section 10.4.3) and point sources (see section 10.4.5).
• Submit the job.
• Visualize your results using ncview.

Very Short-lived Substances

Biogenic emitted Very Short-lived Substances (VSLS) have a short chemical lifetime in the atmosphere compared to tropospheric transport timescales. As the ocean is the main source of the most prominent VSLS, bromoform (CHBr3) and dibromomethane (CH2Br2), this leads to large concentration gradients in the troposphere. The tropospheric depletion of CHBr3 is mainly due to photolysis, whereas for CH2Br2 the loss is dominated by oxidation by the hydroxyl radical.
(OH) both contributing to the atmospheric inorganic bromine (Bry) budget. Once released, active bromine radicals play a significant role in tropospheric as well as stratospheric chemistry as they are in particular involved in ozone destroying catalytic cycles. Although the bromine budget in the stratosphere is dominated by the release of Bry from long-lived source gases (e.g. halons) which is relatively well understood the contribution of biogenic VSLS to stratospheric bromine is still uncertain.

In this exercise the fast upward transport of both VSLS from the lower boundary into the upper troposphere / lower stratosphere (UTLS) due to the super-typhoon Haiyan will be simulated.

In order to perform the simulation, you have to do the following steps:

- Inside the ART-INPUT folder you will find a folder called CASE2-VSLS containing all input data required for this test case.

- Inside $ARTDIR/runctrl/examples/run_scripts you will find the run script for this test case called exp.art.trng19.case2.vsls. Edit the run script according to the namelist parameters you find in Section 10.4. You will find a ? at all places where you have to edit something.

- Create the XML file that is needed to define tracers (see section 10.4.3). For this configuration, you will have to add tracers for CHBr3 and CH2Br2.

- Don’t forget to modify your output namelist accordingly.

- Submit the job.

- Visualize your results using ncview.

---

Volcano Eruption

The eruption of a volcano can have a significant radiative impact as well as an impact on air traffic. In this example, you will simulate the dispersion of volcanic ash particles. Within ICON-ART, you have the choice of treating volcanic ash as monodisperse tracers or with prognostic mass and number (i.e. 2-moment aerosol). In this example, we will make use of the 2-moment description. In order to perform the simulation, you have to do the following steps:

- Inside the ART-INPUT folder you will find a folder called CASE3-VOLC containing all input data required for this test case.

- Inside $ARTDIR/runctrl/examples/run_scripts you will find the run script for this test case called exp.art.trng19.case3.volc. Edit the run script according to the namelist parameters you find in Section 10.4. You will find a ? at all places where you have to edit something. For cart_volcano_file, you can use the file ART-INPUT/CASE2-VOLC/volcano_list_Eyjafjokull.txt.
C.10 Running ICON-ART

- Create the XML file that is needed to define tracers (see section 10.4.3). You also have to create an XML file specifying the aerosol modes in the simulation (see section 10.4.4). For the 2-moment description of volcanic ash, you will need the modes asha, ashb and ashc and the according tracers.

- You can use the namelist switch `lart_diag_out` to obtain diagnostical properties like aerosol optical depth and median diameters.

- Don’t forget to modify your output namelist accordingly.

- Submit the job.

- Visualize your results using `ncview`.

Sea Salt Aerosol

Sea salt aerosol is one of the main contributors to natural atmospheric aerosol. With its high hygroscopicity it is a very efficient cloud condensation nuclei (CCN). Within ICON-ART, sea salt aerosol is described as a log-normally distributed aerosol in three modes with prognostic mass mixing ratios and prognostic number mixing ratios. This simulation includes also a nested domain covering Europe and North Africa with a higher spatial resolution. In order to perform the simulation, you have to do the following steps:

- Inside the `ART-INPUT` folder you will find a folder called `CASE4-SEAS` containing all input data required for this test case.

- Inside `$ARTDIR/runctrl_examples/run_scripts` you will find the run script for this test case called `exp.art.trng19.case4.seas`. Edit the run script according to the namelist parameters you find in section 10.4. You will find a ? at all places where you have to edit something.

- Create the XML file that is needed to define tracers (see section 10.4.3). You also have to create an XML file specifying the aerosol modes in the simulation (see section 10.4.4). For the 2-moment description of sea salt, you will need the modes seasa, seasb and seasc and the according tracers.

- Don’t forget to modify your output namelist accordingly.

- Submit the job.

- Visualize your results using `ncview`.
Bibliography


Index of Namelist Parameters

The following index contains only namelist parameters covered by this tutorial. Please take a look at the document

icon/doc/Namelist_overview.pdf

for a complete list of available namelist parameters for the ICON model. ICON-ART specific namelists are described in the ICON-ART documentation.

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