

v3.0.000

LOTOS-EUROS User Guide

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v3.0.000

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1 Introduction

1.1 What can be found in this guide?

This is the User Guide of the LOTOS-EUROS model. The purpose of this guide is to provide information on how to:

- › obtain a copy of the source code;
- › compile and setup a simulation;
- › run the model
- › add new parts to the code

For information about the physical processes and parameterization used in the model we refer to Manders and team [1].

The User Guide contains the following chapters.

- › Chapter 2 describes the various websites related to the model.
- › How to install the model is described in Chapter 3 on the version control system, and Chapter 4 on the structure of the source files.
- › How to compile and run the model is described in Chapter 5. This requires proper configuration of paths to data and run directories, compiler names, library locations, etc.; details are discussed in Chapter 6 on libraries, Chapter 7 on pre-processing macro's, and Chapter 8 on input data.
- › Chapters 9 on the grid, 14 on the chemistry schemes cbm4 and cbm7, 11 on meteorological data, 13 on the vertical levels, 14 on the chemistry code, and 16 on output are useful when a user would like to adapt the configuration of the model.
- › The final chapters discuss more in-depth elements such as employing parallel computation (18), post-processing tools (19), additional tools (20), and coding conventions (21).

2 Websites

2.1 Public model website

The main source of information on LOTOS-EUROS is the public website:

airqualitymodeling.tno.nl/lotos-euros

The website contains general information, links to the operational forecasts, documents, an overview of publications, and information on how to obtain the open-source version.

2.2 SharePoint site

Documents and data sets specific for the open-source version can be found on the SharePoint site (username/password requested):

<https://365tno.sharepoint.com/teams/T92252/>

3 Version control system

The open-source version of LOTOS-EUROS is not available in an open version control system. Instead, package files are made available to users through the OpenLE SharePoint site (see section [2.2](#)).

Users are encouraged to upload their contributions to the model to the sharepoint site. The development team of LOTOS-EUROS will then take care of integration within new releases.

4 Source files

4.1 Identification of a model version

Each model version is given a three-leveled key, for example "v3.0.000". Here, the first 2 numbers identify the base version number ("v3.0") while the third is a patch number ("000"). New base versions are usually released before the beginning of a new year. Throughout the year, patches might be released to fix a bug or to add new functionality to the current base version.

If you are a new user it is advised to simply use the latest version. As explained in the next section, list the directory 'base' to see what is the latest patch in your copy.

4.2 Directory structure

The LOTOS-EUROS model is shipped in the following directory structure:

```

lotos-euros/v3.0/          # Version directory
README                    # First aid information.
base/                     # Base sources
000/                      # Patch directories
001/                      #
:                          #
proj/                     # Modifications to the base source
tools/                    # scripts for meteo preprocessing etc.

```

At the level of the patch number (000 etc.), a number of subdirectories are present with prescribed names that help to keep Fortran files, scripts, and other data clearly separated from each other:

```

base/000/src/             # Fortran sources
base/000/bin/             # scripts
base/000/rc/              # configuration files
base/000/data/            # data tables
:
proj/testchem/000/src/
proj/testchem/000/bin/
:

```

During the setup of a run, a copy of the source files is created in a temporary build directory, and then compiled over there. The source directories are therefore not polluted by object and module files.

4.3 Patch directories

A patch is an official update of the base, for example with a bug fix or with a new feature that is supposed to become part of a new release. The code of a patch version is stored in the 'base' directory:

```

base/000/                 # initial base version
base/001/                 # first update

```

```
base/002/      # second update
base/003/      # ...
:
```

The complete code of a base+patch is stored in the patch directory.

In the rest of this document we will simply refer to a 'base+patch' code as a 'base' version. If not specified explicitly, the initial patch number is '000'.

4.4 The concept of source projects

A base directory like 'base/000' contains a frozen version of the model. A base version should run without any problem.

In addition to a base, a project could be defined as a modification of files in the base source. A project could also include new files that are not part of the base yet. Typically, projects are defined to:

- › implement a non-standard feature that is very specific to an infrequently used application.
- › create special output required for some applications.
- › test new module features.

For example, a source code could be combined from the following directories:

```
base/000/src /      # base+patch
proj/testchem/000/src / # test code
proj/myoutput/000/src / # user specific
```

In here, the 'base/000/src' directory contains a complete model source, from which some files are replaced by those from the 'proj/testchem/000' directory, and in addition by some from the 'proj/myoutput/000' directory. The order is important: the latest version of a file that is copied will be the one that is actually compiled.

A list of projects to be used should be specified in the run configuration file (explained in section 5.5). For this example it will look like:

```
my.source.dirs      : base/000 \
proj/testchem/000 \
proj/myoutput/000
```

Although not necessary, it is good practice to include the patch number as a sub-directory of a project. In this way it is immediately clear to which patch this project is an extension.

5 Running LOTOS-EUROS

This chapter describes how to configure and start a run with the LOTOS-EUROS model. In section 5.1 a quick start is presented; from section 5.2 onwards the steps are explained in more detail.

5.1 Quick start

LOTOS-EUROS can be run by taking the following steps. This is just to give a first impression or a quick reminder; for details, take a look at the sections that follow.

1. Go to the required version directory:

```
cd lotos-euros/v3.0
```

2. Decide which patch number should be used, for example '000'. Template settings for this patch are available in:

```
base/000/rc/lotos-euros.rc
```

This is the main rc file and once the version is properly installed often the only file that has to be adapted. Browse through it to see if the settings (e.g. run identification, time, domain, species, emissions) for your run are ok. Eventually create a copy and modify for a specific run.

3. Setup a run-directory, compile an executable, and run or submit the job script(s) using:

```
./base/000/bin/setup-le base/000/rc/lotos-euros.rc
```

5.2 Version directory

The best place to setup and start the model is from what we will call the '*version directory*' directory. Change to that directory before following the next steps:

```
cd lotos-euros/v3.0
```

5.3 Run scripts

The script(s) used to setup, start, and submit a LOTOS-EUROS run are placed in:

```
base/000/bin/
```

The 'setup-le' script is the main script for each run.

5.4 Python files

Tools used to setup a simulation are placed in:

```
base/000/py/
```

5.5 Rc-files

The configuration of a model run is done through a text file called the 'rc'-file. The abbreviation 'rc' comes from 'resource', or actually the Unix 'X-resource' file on which the format is based. However, also 'run configuration' is a suitable expansion.

5.5.1 Overview of rc-files

The top-level rc file 'lotos-euros.rc', contains the definition the run id, selects a version the base source and eventually project code, defines a time range and domain, selects emissions and boundary conditions, selects the chemistry scheme, etc. To keep the main file as short as possible, detailed configurations are included from other rc files:

- › lotos-euros-regions.rc in which several model domains are predefined;
- › lotos-euros-landuse.rc in which settings and file names for the land use files are defined;
- › lotos-euros-meteo-*.rc in which paths and settings for input meteorology are defined;
- › lotos-euros-data-meteo-*.rc in which the meteorology model specific data are specified;
- › lotos-euros-radiation.rc in which radiation wavelength bands are defined;
- › lotos-euros-emissions.rc in which paths and settings for anthropogenic emissions are defined;
- › lotos-euros-bound-*.rc files in which paths and settings for several sets of boundary conditions are given;
- › lotos-euros-output.rc in which one can specify the species and fields that have to be put out;
- › lotos-euros-output-label.rc in which the output species and fields for a labelling calculation are defined;
- › lotos-euros-expert.rc in which low-level expert settings are defined, like the chemistry scheme to be used, some model specific settings, IO settings, etc.

Also a number of build/compile/run settings are need; these have been distributed into:

- › lotos-euros-jobtree.rc that defines the jobs that together perform a simulation, e.g. to build, initialize, and run a job.
- › lotos-euros-build.rc in which macros and supported species are defined (section 5.5.6).
- › machine-*.rc files with system-specific settings like compiler and libraries;
- › compiler-*.rc files with compiler specific flags;

The top-level rc file lotos-euros__anywhere.rc contains with default settings for running outside the European domain. It configures a run to use land use and emissions sets that are available globally, but might be less detailed as what has been collected for Europe.

5.5.2 Start: copy from template

To configure your own run, best is create a sub-directory to store your settings:

```
mkdir -p proj/mytest/000/rc
```

This defines that for your project 'mytest' you will use patch '\modelpatch'. The sub-directory 'rc' is used to keep settings separated from source files (that you might want to change too in future).

Then make a copy of one of the template top-level rc-files:

```
cp base/000/rc/lotos-euros.rc proj/mytest/000/rc/lotos-euros.rc
```

Modify it using a text editor, or at least browse through it to see if the settings for your run are ok. It should be self-explanatory with sections for time, grid, species, chemistry, emissions, boundary conditions, landuse, output, use of restart files, etc. Below, the format is explained and some .rc files are treated in more detail. The others should be self-explanatory when opened in an editor.

5.5.3 Format of the rc-file

In this section the ideas and conventions of the .rc file are explained. A simple example of a part in the rc-file could be:

```
! name of input file :
le.input : /data/a.txt
```

This assigns the value "/data/a.txt" to the key "le.input". Tools are available for the run scripts and the model to read values from an rc-file given the keys. The basic format rules for the rc-file are:

- › Keys and values are separated by ':' .
- › Empty lines are ignored.
- › Comment lines start with '!' and are ignored as well.

More advanced usage is possible, for example expansion of keys or environment variables, and conditional setting using if-statements. For a description of the features see the header of:

```
base/000/py/rc.py
```

The best way to learn about all configuration options is to browse through the template rc-file. The added comments explain the use of the various keys and the values they can take.

5.5.4 The machine-specific rc-file

An import setting in the main rc-file is the selection of the 'machine' specific rc-file. This file contains all settings specific for the computer on which the model is running, e.g. compiler settings, library locations, data locations, etc. A template is available:

```
base/000/rc/machine-template.rc
```

For each institute/machine combination, a machine-specific rc-file should be created. The settings are included in the top-level rc-file via:

```
! include settings:
#include base/${my.patch.nr}/rc/${my.machine.rc}
```

When GNU environment modules are used on your computer, the machine settings might be used to insert 'module' commands in the top of job scripts. For the TNO server this is for example used to load the correct environment (using gcc compiler version 13.1.0):

```
! modules loaded in job scripts:
*.modules : purge ; \
use /tsn.tno.nl/Data/SV/sv-059025_unix/admin/${SERVER}/modulefiles ...
...; \
load slurm ; \
load ${my.compiler.suite}-suite/${my.compiler.version} ; \
load openmpi/5.0.3 ; \
load hdf5/1.14.4-2 ; \
load netcdf-c/4.9.2 ; \
load netcdf-fortran/4.6.1 ; \
load udunits/2.2.28 ; \
load lapack/3.12.0 ; \
load spblas/1.02.917 ; \
load nco/5.2.4 ; \
load makedepf90 ; \
load lotos-euros
```

See chapter 6 for information about the external libraries that should be configured in the machine rc file.

5.5.5 The compiler rc-file

The machine-specific rc-file includes a file with compiler-specific settings:

```
! GCC compiler suite, selected with environment variables:
#include base/${my.le.patch}/rc/compiler-${my.compiler.suite}-${my...
...compiler.version}.rc
```

For each compiler suite used to compile the model, a compiler-specific file should be present.

5.5.6 The build rc-file

Many settings that have to do with the setup and installation of the model have been hidden for the users by collecting them in the 'build' rc-file, which is included into the main rc file:

```
! include expert settings to build source code
#include base/${my.le.patch}/rc/lotos-euros-expert.rc
```

Do not modify this file unless you know what you are doing!

5.6 Setup script

To setup a run, first go to the model version directory:

```
cd lotos-euros/v3.0
```

Call the *setup* script with the rc file as argument to create a run-directory and compile an executable:

```
./base/000/bin/setup-le proj/mytest/000/rc/lotos-euros.rc
```

To see the extra options that are accepted by the setup script, use:

```
./base/000/bin/setup-le --help
```

Note that all 'long' options like '--help' usually have a 'short' version too, in this case '-h'.

A useful option is '--new' or '-n', which first removes the existing build directory, followed by creating a completely new one. In case you receive strange errors from the compiler, that might have to do with messing up old and new objects, so try whether this option solves the problems.

5.7 The job-tree

The 'setup-le' script will start a sequence of jobs to initialize and run a LOTOS-EUROS simulation.

By default the jobs are:

- › le.copy job will copy the source codes to a run directory;
- › le.build job will compile an executable;
- › le.run job will run the executable, this is the actual simulation;
- › le.post job will post-process the output (if necessary).

The list of jobs to be created is defined in:

```
lotos-euros-jobtree.rc
```

with:

```
jobtree.le.elements : copy build run post
```

The 'le.copy' job will always run in the foreground.

The 'le.build' job should probably run in foreground too, since this helps to quickly see compilation errors. This is defined in:

```
lotos-euros.rc
```

with:

```
! copy job is running in foreground (see "lotos-euros-jobtree.rc"),
! enable the following line to have also the build job in ...
...foreground:
jobtree.le.build.script.class : utopya.UtopyaJobScriptForeground
```

The default destination for the 'run' and other jobs is probably to submit it to a queue system. Since the queue system is different on each platform, the destination is defined in the machine-specific settings (section 5.5.4). For the TNO platform the SLURM queue should be used:

```
! default class with the job script creator:
*.script.class : utopya.UtopyaJobScriptBatchSlurm
```


This defines that the header of a job file should have comments with SLURM job options. See the examples of machine specific settings for inspiration on configure job settings for your own queue system.

5.8 The 'le.copy' job

This job is the first that is performed and will create a run directory with a copy of the source code.

5.8.1 Run directory

The first step taken by the 'le.copy' job is to create a run directory on a location specified in the rc-file. Typically the run-directory will be located on a scratch space, since a lot of temporary files are created while running that do not have to be backed-up. A typical content of the the run directory is:

```
<rundir>/build/      # source code, object files
<rundir>/run/        # executable, rc-files, submit script, log files
<rundir>/output/     # output files
<rundir>/restart/    # restart files }
```

5.8.2 Run-time rc-file

The 'copy' job will also create a *run-time* rc-file. This a single settings file with all includes and variable substitutions resolved; see section 5.5.3 for details.

The run-time rc-file is put in the directory:

```
<rundir>/run/lotos-euros.rc
```

5.8.3 Collection of a source code

The next step is the collection of a source code in a build directory. The build directory is usually located on a temporary scratch disk, the exact location is specified in the rc-file. The sub-directory where the code is collected includes the name of the compiler and the choices for the compiler flags (see section 5.9.3 for the compiler flag settings). This is done to ensure that an executable is compiled with the same flags applied for all source files. An example of a sub-directory name:

```
<rundir>/build_optim-none_check-all/src/
```

The concept of source projects is explained in section 4.4. In summary, the first collected source files are those in the 'base/xxx' directory. Subsequently, the files from the base version are overwritten by patches or project specific versions from 'proj/' directories according to the specification in the main .rc file.

For example, the rc-file might contain the following setting for the list of source directories to be included:

```
my.source.dirs      : base/000 \
proj/testchem/000 \
proj/myoutput/000
```

In this example, the LOTOS-EUROS source is formed from:

1. the files in 'base/000/src' ...
2. ...replaced with equally named files from 'proj/testchem/000/src/' ...
3. ...replaced with equally named files from 'proj/myoutput/000/src/'.

The test files in 'proj/testchem/000/src/' will not affect the 'official' versions of the code. Thus, if you want to change something in the code, create a new project directory and implement your changes there.

5.9 The 'le.build' job

The 'le.build' job configures the source files and compiles an executable.

5.9.1 Generation of source files

Some source code files are generated by the scripts using settings in the rcfile and data tables. These are:

- › source files with tracer definitions and chemistry codes (see section 14);
- › preprocessing macro include files (section 7).

Default versions of the generated source files are included in the base source. Editing won't have any effect; instead change the rcfile settings or the tables that were used to generate them.

5.9.2 Creation of the Makefiles

The source directory of LOTOS-EUROS contains makefile dependencies in:

Makefile_deps

This file has been created using the 'makedepf90' program, and should be sufficient for the standard source. In case dependencies are changed, for example if new source files are added, either edit the dependencies file (might be difficult), or ensure that 'makedepf90' is available on your system. The program can be downloaded from the the MakeDepF90 website¹. To install it under your own account, use something like:

```
tar x -f makedepf90-2.8.8.tar.gz
cd makedepf90/2.8.8
configure --prefix=${HOME}/opt/makedepf90/2.8.8
make
make install
```

and ensure that the paths are set:

```
export PATH="${HOME}/opt/makedepf90/2.8.8/bin:${PATH}"
export MANPATH="${HOME}/opt/makedepf90/2.8.8/man:${MANPATH}"
```

When installed (or already present), change the following flag in the machine rcfile (section 5.5.4):

¹<http://personal.inet.fi/private/erikedelmann/makedepf90>

```
! Is makedepf90 installed?
my.with.makedep          : True
```

This flag is then used in the expert settings (section 5.5.6) to re-create the makefiles automatically by a call to the 'makedepf90' program.

5.9.3 Compilation

The final step performed by the 'setup-le' script is the compilation of the executable.

An important configuration choice for compiling is setting the compiler flags. By default, the executable is compiled with the 'fast' flags to have a run-time as low as possible. For testing and debugging it is however useful to enable the 'check' flags, which could for example trap out-of-array-bound problems and floating-point-exceptions (division by zero etc). *After changing the code, first run with checks enabled!*

The flags to be applied can be set in the pycasso rc file by a list of keywords. The default setting for compilation with the fast flags is:

```
my.build.configure.flags      : optim-fast
```

For testing and debugging purposes, the 'check' flags can be turned on with:

```
my.build.configure.flags      : optim-debug check-all debug
```

The actual flags assigned to these keywords are set in the compiler rc-file described in section 5.5.5.

5.10 The 'le.run' job

This is the actual run.

The executable is started either directly, or via an MPI started to have the correct environment for the domain decomposition.

5.11 The 'le.post' job

This job is currently only used to compress output files. The compression is done using the 'NCO' tools. Often, output data files contain many similar values (zeroes or empty fields). In these cases, compression of those files leads to considerably smaller files without loss of data.

6 External libraries

6.1 Library settings

The location of external libraries such as NetCDF is strongly dependend on the machine where the model is compiled. Configuration is therefore done in the machine settings file described in section 5.5.4.

6.2 Librararies used in base code

The following external libraries are currently used, of which NetCDF is the only obligatory.

6.2.1 NetCDF library

This library is required for:

- › reading meteorological and other input data;
- › writing model output.

The model does not use NetCDF-4 feautures yet; it is therefore sufficient to link with a NetCDF-4 library that was compiled without NetCDF-4 features enabled, or even with a classic NetCDF-3 library.

The machine-specific settings (section 5.5.4) specify the compile and link flags for this library, for example:

```
compiler.lib.netcdf.fflags : -I/opt/include
compiler.lib.netcdf.libs   : -L/opt/lib -lnetcdff -lnetcdf
```

Note that in case shared libraries are used (as in this example), then it might turn out to be necessary to tell the linker to add the path to the shared libraries to the runtime search path:

```
compiler.lib.netcdf.libs : -L/opt/lib -lnetcdff -lnetcdf \
-Wl,-rpath -Wl,/opt/lib
```

6.2.2 UDUnits library (optional)

The UDUNITS library is used to check if units read from a file match with the units used in the model. Some common unit comparisons have been hard-coded, so that for example for the test input provided with the open-source version the UDUNITS library is actually not necessary. In that case, the need for a UDUNITS library could be disabled in the machine rcfile (section 5.5.4) using the empty setting:

```
my.udunits.define :
```

If the model complains however that a certain unit check cannot be performed, either hard-code a comparison on the location identified by the error message, or enable the need for a UDUNITS library. The UDUnits library might be available as version 1 or 2; since the interface

is rather different between these, it was necessary to distinguish between them in the code. Therefore, the flag in the machine settings should define explicitly which version is available on the system:

```
!~ version 1 is available:
!my.udunits.define      : with_udunits1
!~ version 2 is available:
my.udunits.define       : with_udunits2
```

Note that for version 2 a C-binding is used that requires a compiler with F2003 support.

6.2.3 Pandas Python library (optional)

The Python scripts that generate chemistry source files for the model (GENES scripts, described in Chapter 14) use the Pandas library when the CBM7 chemistry scheme is selected. This has been tested with Pandas version 2.1.4, older versions are likely to work as well.

6.3 Adding new libraries

To compile the model with new libraries, configurations are required at a number of levels.

- › The user should explicitly enable the required libraries using macro definitions. For example, to compile with the HDF5 library, introduce a new macro 'with_hdf5' following the instructions in chapter 7.
- › Ensure that this macro is defined by adding it to the 'define' list in the expert settings (section 5.5.6):

```
build.configure.macro.define : with_netcdf with_hdf5
```

- › For this macro, provide a list with library names that should be linked with. For example, if HDF5 is required, then also a number of compression libraries should be linked. For this, the expert settings should contain a definition similar to:

```
build.configure.libs.ifdef.with_hdf5 : hdf5 sz jpeg z
```

- › The order in which the libraries should be linked is defined by:

```
build.configure.libs.all : netcdf hdf5 sz jpeg z
```

- › The machine-specific settings (section 5.5.4) specify the compile and link flags for a library, for example:

```
compiler.lib.hdf5.fflags : -I/opt/include
compiler.lib.hdf5.libs   : -L/opt/lib -lhdf5_hl \
-lhdf5_fortran -lhdf5
```

7 Pre-processor macro's

Preprocessing macro's are a convenient tool in programming large applications. This chapter describes the use of these macro's in the LOTOS-EUROS source.

7.1 Expert settings

Working with macro's is considered an expert job, since small typo error could cause a part of the code to be omitted without noticing. Therefore, the expert rcfile includes lists of macro's that are supported. The source code is checked on use of macro's that are not supported; if these are found, an error is raised and configuration stops. The lists in the expert rcfile are used to add macro definitions to the proper macro include file.

7.2 Example of usage

Sometimes a minor modification of the actual source code is needed, for example because certain compilers cannot digest a piece of code, or to enable code that depends on an external library that might not be available (but is not always needed either). For this so-called pre-processing macro's are used. For example, the following code hides the use of the HDF4 library that is usually only needed for very specific input files:

```
#ifdef with_hdf4
! open hdf file :
call HF90_Open( 'input.hdf', HF90_READ, hdf_id, status )
...
#else
stop 'not compiled with HDF4 library enabled'
#endif
```

In this example, if the macro 'with_hdf4' is defined, then only the code between '#ifdef' and '#else' is used, otherwise the code between '#else' and '#endif' is used. If the code is compiled with the HDF4 library enabled, then the macro 'with_hdf4' should be defined and the file 'input.hdf' will be opened as specified, if this piece of code is called. However, a user wants to run the model, but does not need to read HDF4 files, the macro definition could be omitted. This is in particular useful if this library is not available; the user should not be bothered with a compiler complaining that a library is not available while it is not used anyway.

7.3 Macro definition include files

Macro's are defined by statements like:

```
#define with_hdf4
```

All macro definitions are collected in small include files. For example, for macro definitions in LOTOS-EUROS source files the include file 'le.inc' could look like:

```
!
! Include file with macro definitions.
!
```

```
#define with_hdf4
```

This file is included in the header of a file with:

```
#include "le.inc"
```

7.4 User settings

Which macro's are defined is something that is often user and application dependent. Therefore, a list of macro's to be defined is part of the rcfile:

```
my.le.define    : with_hdf4
```

From the values in this list the macro include file(s) are written automatically by the source configuration scripts.

8 Input data

8.1 Test package

A package with input data is available to run the model with the default settings to verify proper installation. It can be downloaded from the TNO FTP site (web-ftp81.tno.nl).

Specific input files need to be retrieved separately. For instance, MACC III emission data are not included but can be obtained upon request. On the other hand, meteorological data and boundary conditions from MACC cannot be provided by TNO.

8.1.1 Content

The content is as small as possible, but large enough for a model run with the following properties:

- › simulation period August 2012;
- › European domain as used for operational forecast;
- › MACC-II emissions 2009);
- › boundary conditions from climatology

The directory tree of the data is as follows:

```
inputdata /
  /ammonium/
  /bound/
  /cf-standard /
  /emissions/
  /CAMS/REG/v2_2_1/
  /LEIP/
  /europe_w30e70s5n75/
  /ECMWF/od/ # meteo
  /MACC/ fire /
  /landuse/
  /forest/
  /soiltext/
  /soilwater_avg/
  /traffic/
  /standard/
```

8.1.2 Settings

The location of the input data is machine specific and therefore set in the 'machine.rc' file:

```
my.data.dir      : /data/inputdata
```


8.2 See also

For more detailed information on certain types of input data, see:

- › chapter 11 on meteorological data;
- › chapter 12 on boundary conditions.

9 Horizontal grid

9.1 Grid definition

Two different grid types are supported: a regular longitude/latitude grid ("cartesian"), and a grid that is curved in longitude/latitude sense ("non-cartesian").

9.1.1 Cartesian grid

The default grid for most runs is regular in longitudes and latitudes, thus with equal spacing in degrees in both directions. In longitudes/latitudes the domain is square (figure 9.1, left panel).

The rcfile should first specify the grid type:

```
! define grid type:
grid.type          : cartesian
```

For this grid type, the settings should then specify the lower-left corner, the resolution, and the grid size:

```
grid.west          : -15.0
grid.south         : 35.0
grid.dlon          : 0.50
grid.dlat          : 0.25
grid.nx            : 100
grid.ny            : 140
```

Template grid definitions are available in:

lotos-euros-regions.rc

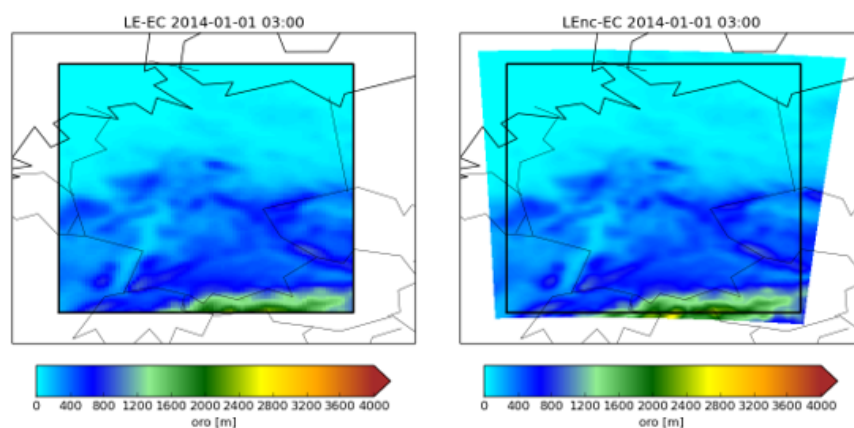


Figure 9.1: Examples of model extend on cartesian grid (left) and non-cartesian grid (right).

If this file is included in the main rcfile, the grid definition to be used should be selected using a keyword. For example, the following settings will enable the 'MACC-II' grid definition as used in the operational CAMS forecasts:

```
! select grid used by operational forecasts:
grid.name          : MACC-II

! grid definitions:
#include base/${my.patch.nr}/rc/lotos-euros-regions.rc
```

9.1.2 Non-cartesian grid

A non-cartesian grid is typically used when the model is driven by meteorological data from WRF or COSMO. In that case it makes sense to use the same grid definition as the meteorological model. In longitudes/latitudes the domain is usually wider towards the poles to have grid cells that have equal width in km (figure 9.1, right panel).

In the rcfile, first specify the grid type:

```
! define grid type:
grid.type          : non-cartesian
```

The grid definition should then be read from a sample file with the WRF or COSMO meteo. The settings should specify the sample file name and the name of a variable from which the grid definition will be read, for example:

```
! sample file:
grid.file.name     : /data/WRF/wrfout_d02_2014-01-01_00:00:00
! sample variable:
grid.file.var      : var_name=HGT
```

In practice it is often only necessary to run the model on a sub domain of the input data. The following setting could be used to select this sub domain, or to specify that the full grid should be read:

```
! subset [i1,i2,j1,j2], negatives for all :
!grid.file.subset      : -999 -999 -999 -999
grid.file.subset       : 50 110 50 150
```

9.2 Zooming

Running in zoom modes simply means running the model twice: first for a large domain and coarse resolution, then for a smaller domain in fine resolution. The later should be configured to read concentrations from the first run as boundary conditions.

The following steps are usually taken to setup a zoom run.

1. Perform a run on large domain:
 - › large domain covering the future zoom grid;
 - › coarse resolution;
 - › output of 3D concentration fields:
 - all relevant tracers; use keyword: 'all-advected'
 - bounding box around zoom region to save disk space (section 16.2.1);

- conc-bound output; conc-bound output generate concentrations + pressure and height information of grid cells
- › remember the run-id and the output directory.
- 2. Perform the zoom run:
 - › small domain within the previous domain;
 - › fine resolution;
 - › boundary condition; in main .rc file: Load bound-rc file including settings to use LE-run as boundaries. (lotos-euros-bound-*-le.rc). 'conc-bound' output will be interpolated to zoom-domain
 - › parent domain; Domain name of the boundary run
 - › eventually put out concentrations in the halo cells to check the inheritance of the boundary conditions, see section [16.2.1](#).

Note that the time steps for the zoom run are automatically reduced to match the CFL (Courant)-criterion.

In the setting of the rc-file the type of run can be given ('my.run.type'), the necessary outputs will be defined by default.

- › Bound: run will be used as boundary run; keyword 'all-advected' is used to get all advected tracers in output (conc-bound files).
- › Zoom: run will be used as zoom run; Parent domain should be defined, rc-file including LE-settings will be loaded.
- › Zoom or Bound_plus_Zoom; Run is used as both boundary and zoom run (middle domain in a three-step nested approach).

9.3 Mapping of input data

Input is interpolated or averaged automatically to the required grid; if the model domain extends beyond the coverage of the input, an error is raised.

10 Vertical levels

Three different vertical level definitions are supported. Table 10.2 shows selected properties for the different definitions discussed below.

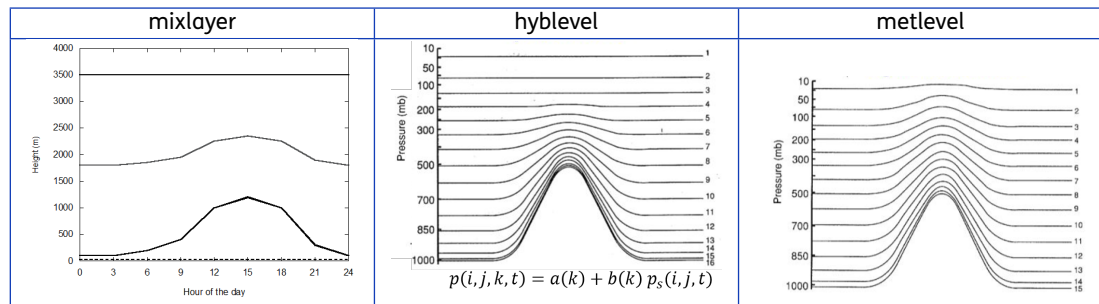


Table 10.1: Illustration of level definitions.

levels.type	mixlayer	hyblevel, metlevel
number of layers	4 or more	unrestricted
vdiff.kz_type	msp (default) normal (optional)	normal
adjust process	yes	no
volumes	constant within time step (1 hour)	dynamically

Table 10.2: Properties of vertical level definitions.

10.1 Mixed-layer definition

The 'mixed-layer' definition was the standard method for many years for the vertical structure in LOTOS-EUROS. The levels are defined every time step in terms of heights above the surface, and some of them evolve dynamically with the boundary layer height obtained from the meteorology (figure 10.1, left panel). Input data will be mapped to this layer definition.

In a standard configuration with 5 layers, the first layer is thin surface layer of 25 m. On this the second layer is placed with the top at the boundary layer; this is therefore grid cell and time depended. The third and forth layer are reservoir layers of equal thickness of at least 500 m, with the forth layer having a top at 3500 m above the surface or more if needed to have the minimal thickness. On top of this the fifth layer is placed with a top at 5000 m or more if needed to have a minimum thickness of 500 m. The minimum thickness is increased with a factor times the orography of surrounding cells to have somewhat thicker cells over mountain area's; by using thicker cells, the minimum time step implied by the advection is higher which saves run time.

This level type is enabled using:

```
levels.type           : mixlayer
```

The number of layers to be used is defined with:

```
levels.nz             : 5
```

The thickness of the surface layer is defined with:

```
mixlayer.surf_top      : 25.0
```

The mixing layer should be at least as thick as the surface layer, which is defined by a minimum value for the top above the surface:

```
mixlayer.mix_topmin   : 50.0
```

The fifth and higher layers are defined using lists that define the top above the surface, the minimum thickness, and the factor to be used in combination with the standard deviation of the orography. For the 5 layer version the configuration is:

```
mixlayer.top          : 3500.0 5000.0
mixlayer.dmin         : 500.0
mixlayer.sdofac       : 1.0 0.5
```

For an 8 layer version the configuration looks like:

```
mixlayer.top          : 3500.0 5000.0 6500.0 8000.0 10000.0
mixlayer.dmin         : 500.0
mixlayer.sdofac       : 1.0 0.5 0.0 0.0 0.0
```

A final configuration is the definition of the thickness of the aloft layer that will hold the top-boundary concentrations:

```
mixlayer.daloft       : 1000.0
```

With this vertical scheme, one of the model operators is an 'adjust' step that changes the layer heights towards a new boundary layer height. This is applied at the start of every time step. During the time step, the grid cell heights (and thus cell volumes) are kept constant.

For diffusion between two layers two options are available to compute diffusion coefficients K_z at the layer interfaces. The first is the 'msp' method that has been used traditionally, which adjusts the K_z values for differences in layer thickness around the interface; this is enabled with:

```
vdiff.kz_type : msp
```

Alternatively also the 'normal' method could be enabled that is used by the other level definitions.

10.2 Hybrid layer definition

The 'hybrid layer' definition is the current standard method for the vertical structure in LOTOS-EUROS. A hybrid-sigma-pressure scheme defines vertical layers using pressure boundaries that follow the surface pressure at the lower levels, and have fixed pressure boundaries at the top. The layers therefore follow the orography near the surface, which evolves slowly to layers at fixed pressure higher up (figure 10.1, middle panel). This scheme is for example used by the ECMWF meteorological model, and is therefore useful if LOTOS-EUROS is driven by this data. Input data will be mapped to the chosen layer definition anyway, regardless whether this is also a (different) hybrid definition, or something else.

At every time step, the pressures at a layer interface are

$$p(i, j, k, t) = a(k) + b(k) p_s(i, j, t) \quad (10.1)$$

In here, (i, j) denotes the horizontal cell indices, k the layer interface index, and t the time. The interface coefficients a and b define for each layer which part fraction of the pressure is constant and which fraction depends on the surface pressure p_s that is part of the meteorological input.

The hybrid level definition is enabled using:

```
levels.type : hybrlevel
```

The number of layers to be used is then defined with:

```
levels.nz : 12
```

The rest of the atmosphere (above the layers simulated by the model) is filled with concentrations from the global boundary conditions; these are used for top boundary conditions, and to simulate satellite observations. Specify the total number of layers with:

```
levels.nz_top : 18
```

NOTE: the first top layer above the model layers should not be too thick! This will be filled with concentrations from the global boundary conditions, averaged over the layer. If the layer is too thick, the ozone concentrations will include part of the stratospheric ozone layer, and the inflow of ozone from the top will be too high!

The hybrid coefficients a and b should be provided in a text file, for which an example is provided with the model distribution:

```
hybrlevel.coefficients : data/hybrlevel-137_CL42_ml12p06
```

The number of coefficients in the file might exceed the requested number; the model will simply read the first values that define the interfaces from surface to model top.

In this scheme, the layers are first defined in terms of pressure boundaries, which then implies the heights above the surface (using temperature, humidity, and orography). The cell volumes change dynamically but do not follow the evolution of the boundary layer, and therefore no 'adjust' process is needed as used for the 'mixlayer' scheme.

Diffusion coefficients between layer interfaces should be normal K_z values, thus not adjusted for extreme differences in layer thickness:

```
vdiff.kz_type : normal
```

10.3 Meteo level definition

With this method, the model will copy a level definition from a set of meteorological data. Depending on the data this definition could define layer interfaces as pressures or heights above the surface. The model data settings as described in chapter 11 should define the correct settings to read and process the data. This method is useful to keep LOTOS-EUROS as close to a meteorological model as possible. The 'meteo-level' definition is therefore a generalization of the 'hybrid-level' method, which does not necessarily require a surface pressure and hybrid coefficients but could use other level definitions too (figure 10.1, right panel).

The meteo level definition is enabled using:

```
levels.type      : metlevel
```

The layers to be used are defined with two settings: the number of layers, and for each of these, the number of original layers that is coarsed in them:

```
! number of layers that should be used:
levels.nz      : 12
! layer combination, empty for no combination:
metlevel.combine : 1 1 1 2 2 2 2 2 2 2 2 2
```

The rest of the atmosphere (above the layers simulated by the model) is filled with concentrations from the global boundary conditions; these are used for top boundary conditions, and to simulate satellite observations. Specify the total number of layers, and how to coarsen them from the meteo layers using:

```
! total number of layers including "top"
levels.nz_top      : 19
! layer combination, empty for no combination:
metlevel.combine_top : 1 1 1 2 2 2 2 2 2 2 2 2 1 2 3 3 4 4 4
```

NOTE: the first top layer above the model layers should not be too thick! This will be filled with concentrations from the global boundary conditions, averaged over the layer. If the layer is too thick, the ozon concentrations will include part of the stratospheric ozone layer, and the inflow of ozone from the top will be too high!. So first top-layer should not be a combination of layers but consist of only 1 single layer

Using a layer definition from a meteo model requires different settings depending on the originating model. An example of such a specific configuration is provided as:

```
rc / lotos-euros-data-meteo-cosmo-metlevel.rc
```


11 Meteorological data

11.1 Introduction

The LOTOS-EUROS model uses off-line meteorology. Meteorological fields are read from files with time series of data at for example 3 hourly resolution.

The storage and reading of meteorological fields has been revised completely for version 2.0. The new implementation is based on general routines that are able to handle data files in NetCDF format following common conventions.

Previous versions of the model also supported meteorological data from the RACMO regional climate model and the WRF meteorological model. The new generic interface of the model will be extended to support data files produced by these models too.

11.2 Data definition in model

The meteorological data in LOTOS-EUROS is allocated dynamically, and only if actually needed. The dynamic allocation is part of generic facility in the model that is intended to hold all gridded variables, but as first step holds the meteorological variables.

Definition of the meteorological variables is done in the settings file(s), in particular:

`lotos-euros-data.rc`

The header of the files contains details of the possible settings, here we describe the main steps only. The first definition is a (long) list of all data variables supported by the model. This list contains for example a keyword to describe the 2m surface temperature:

```
data.vars      :    ... tsurf ...
```

For each of the variables, a detailed description should be provided in the form of specific settings lines. For the surface temperature 'tsurf' this is for example:

```
! define :
data.tsurf.long_name      : surface temperature
data.tsurf.units          : K
data.tsurf.range          : 0.0 Inf
data.tsurf.gridtype       : cell
data.tsurf.levtype        : sfc
data.tsurf.datatype       : instant_field_series
data.tsurf.input          : meteo.tsurf
```

The detailed settings contain (see also the header of the settings file):

- › a descriptive *longname*, used when the variable is put out;
- › the units;
- › the range of allowed values, used to truncate the variable to realistic values (sometime necessary to remove tiny negative values for example);

- › the horizontal grid on which the variable is defined;
- › the vertical levels on which the variable is defined;
- › the datatype (Fortran class), which is mainly related to the temporal behavior (see section 11.6);
- › a keyword that describes the input mechanism in case the variable should be read from file(s) (see section 11.7); otherwise, a definition of how to compute the field should be present (see section 11.8).

11.3 Accessing a variable in the model

In the model code, the variables that are defined in the settings are available from a module:

```
use LE_Data
```

The array that holds the variable values should be accessed using a pointer. Use the following code to access the above defined surface temperature:

```
use LE_Data only : LE_Data_GetPointer

! this will point to the surface temperature:
real, pointer          :: tsurf(:, :, :) ! (lon, lat, 1)

! assign pointer to surface temperature,
! check if the units are as expected:
call LE_Data_GetPointer( 'tsurf', tsurf, status, check_units = 'K')
IF_NOTOK_RETURN(status=1)

! show value in cell (3,4) :
print *, 'example of surface temperature: ', tsurf(3,4,1)
```

Note that:

- › the variables are always 3D, for surface fields the last dimension has size 1;
- › a check on the units can be used to make sure that the units are as expected (thus degrees Kelvin and not degrees Celcius);
- › the call to the 'GetPointer' routine will return with an error status if the requested variable was not defined in the settings, or when it was not actually enabled (see section 11.4).

11.4 Enabling a variable in the model

Before a variable could be used, it should not only be defined in the settings, but also actually enabled. By enabling a variable it will be allocated and filled with the proper values at every time step.

Enabling is typically done in the initialization routine of a module, where all variables should be enabled that are used in the module. If a variable is not enabled, for example because some processes in the model are not used in a particular configuration, it will not be allocated (which saves memory) and will not be read or computed (saves run time).

To enable a variable, use the following lines of code:

```
use LE_Data, only : LE_Data_Enable

! enable surface temperature:
call LE_Data_Enable( 'tsurf', status )
IF_NOTOK_RETURN(status=1)
```

11.5 Grid types

The following grid types are supported:

- › **cell**
Values valid for the entire cell; this is the most common type. The shape of this grid is '(nlon,nlat)'.
- › **corner**
Values are defined on the corners of cells. This is used to setup the advective fluxes through the cell edges. The shape of this grid is '(nlon+1,nlat+1)'.
- › **u-edge, v-edge**
Used for fluxes through the west and east sides ('u'), or through the north and south sides ('v'). The shape of this grid is '(nlon+1,nlat)' or '(nlon,nlat+1)' respectively.

When fields are read from input files (section 11.7), a re-mapping to the target grid is performed automatically if possible. Current implementations can handle longitude-latitude grids (eventually with irregular spacing), and to some extent the so-called reduced grids which have a varying number of cells per latitude band.

11.6 Data types

The variable definition contains a setting for the datatype:

```
data.tsurf.datatype          : instant_field_series
```

The following data types are possible:

- › **field**
This type is used for a constant field, thus without any change in time, for example the grid cell area.
- › **instant_field**
An instant field is valid for specific instant time. This type is typically used for fields that are computed from other instant fields, thus not read from a time series in input files.
- › **instant_field_series**
Use this type for a variable that is derived from a time series, typically by interpolation in time between the fields in the series. For example the 2m surface temperature is of this type.
- › **constant_field**
In this context a 'constant' field means 'constant during a time interval', for example during 3 hours. A time interval for which the variable is valid is associated with it. This type is typically used for fields that are computed from other constant fields, thus not read directly from file.

constant_field_series

Use this type to read 'constant' fields from a time series in a file. For example the amount of precipitation is of this type, since in meteorological data is available as the total amount of water that has fallen down during some time interval.

11.7 Input descriptions

If a variable should be read from input files, a configuration of the following form should be present:

```
data.tsurf.input          : meteo.tsurf
```

This setting describes that the details of how the field should be read are defined with settings starting with 'meteo.tsurf'.

For the current model version, meteo files were created from ECMWF data, see section 20.1 for creation scripts. Examples of input descriptions for these files are found in the settings file:

```
lotos-euros-meteo-ecmwf.rc
```

Here we show the general idea of the settings; for the specific settings we refer to the actual rc file.

The input description should first contain an 'input' value that defines time intervals and descriptions:

```
meteo.tsurf.input : 2012-01-01 00:00, 2012-12-31 23:59, fmt1.tsurf | \
2013-01-01 00:00, 2020-12-31 23:59, fmt2.tsurf
```

That is, a list of time intervals and description keys separated by a vertical line ('|') should be provided. Given a requested time, the input will be read according to the description associated with the interval that encloses the time value. This feature is used to handle changes in for example file formats and resolutions over time, which often occur in operational meteo models.

The description key refers to settings that are used to define the file and variable name. For example for the surface temperature these settings could have the form:

```
fmt1.tsurf.name : /data/ECMWF/sfc/{yyyy}/t2m_{yyyymmdd}_3h.nc
fmt1.tsurf.var  : long_name=2 metre temperature;var_name=t2m
```

The 'name' setting provides a template for the file name in which the variable should be found; the keywords '{yyyy}' etc. are replaced by actual time values. The 'var' setting identifies the file variable, and consists of a ';' separate list of 'key=value' pairs. The 'key' could be 'long_name' or 'standard_name' in this case it refers to a variable attribute in the netcdf file; a 'var_name' refers to the variable name itself. A 'standard_name' is preferred since this might be less subject to changes than a long name or the variable name. The first 'key=value' pair that identifies a variable in the file is used.

For instant fields (valid for a single time step), also the temporal interpolation method should be defined. Here we define that the expected temporal resolution in the files is 3-hourly, and that that requested fields should be interpolated linearly in time:

```
meteo.tsurf.tinterp      : interpolation=linear;step=3;units=hour
```

11.8 Computing variables

If a meteo variable should not be read from input files but computed from other variables, the definition should have a 'call' description. For example, the following description is used to define a variable 'srh' with relative humidity at the surface, which should be computed from the 2m surface temperature ('tsurf') and the 2m dewpoint temperature ('dsurf'):

```
data.srh.long_name      : surface relative humidity
data.srh.units          : %
data.srh.range          : 0.0 100.0
data.srh.gridtype       : cell
data.srh.levtype        : sfc
data.srh.datatype       : instant_field
data.srh.call           : RelativeHumidityTD( tsurf , dsurf )
```

The 'call' description has the form of a function call, with as arguments a comma-separated list of source variable names. The function name ('RelativeHumidityTD') is linked to an actual function call in subroutine:

```
Variables_Setup      (module LE_Data_Variables)
```

The arguments of the actual function are usually the name of the target variable ('srh') and a list with the names of the argument variables ('tsurf','dsurf').

If a variable with a 'call' definition is initialized, it is checked if all arguments are names of variables. When the variable is enabled, the arguments are enabled too.

11.9 Example: 3D field

The following example show the definition of a 3D field, in this case temperature. Such a variable combines most of features described in this chapter.

The input time series of temperature is here assumed to have a 3 hourly resolution, and defined on pressure levels. At every time step the temperature on model levels should be computed from a temporal interpolation between the time steps in the input, and a vertical remapping to the model levels (horizontal re-mapping is done at reading from input as described in section 11.5). For the vertical re-mapping it is necessary to have the level definition of the input fields and the model; here we apply an air-mass weighted re-mapping that requires level definitions in term of half-level pressures.

The name of the temperature variable in the model will be 't'. This will be computed from a variable 't_met' which holds the temperature field at the levels of the meteorological model (as read from the input). For the re-mapping also half-level pressures for the model and the meteorological input are needed. Therefore, in total 4 variables are needed in the model:

```
data.vars              : ... hp_met t_met hp t ...
```

The temperature field is defined with:

```
data.t.long_name       : temperature
data.t.units           : K
data.t.range           : 0.0 Inf
data.t.gridtype        : cell
data.t.levtype         : levels
```

```
data.t.datatype      : instant_field
data.t.call          : MassAverage( hp_met, t_met, hp )
```

The variable is defined on grid cells and model levels, and is an instant field valid for a single time step. At every time step, the values should be computed from a mass-weighted average from the temperature at the meteorological levels, which also requires half-level pressure fields.

The temperature at meteorological levels is defined with:

```
data.t_met.long_name      : temperature
data.t_met.units          : K
data.t_met.range          : 0.0 Inf
data.t_met.gridtype       : cell
data.t_met.levtype        : meteo_levels
data.t_met.datatype       : instant_field_series
data.t_met.input          : meteo.t
```

The data type is that of an instant series, since the field should be interpolated in time from a time series of input fields. The input is described by rcfile keys that start with 'meteo.t'. The level type 'meteo_levels' forces the model to maintain the original levels as read from the input files.

The input description of the temperature fields contains a single file format for all times, and tells the model that the input time resolution is 3-hourly and that linear interpolation should be applied in between:

```
meteo.t.input    : 2012-01-01 00:00, 2020-01-01 00:00, ecfile.t
meteo.t.tinterp  : interpolation=linear;step=3;units=hour
```

The file description looks like (see 'lotos-euros-meteo-ecwmwf.rc' for the exact form):

```
ecfile.t.name    : /data/ml-tropo20/t_%{yyyymmdd}_3h.nc
ecfile.t.var     : long_name=Temperature
```

For the definition of the half-level pressure fields that are used for the mass-averaging we refer to the settings in 'lotos-euros-data.rc'.

12 Boundary condition data

12.1 Introduction

The boundary conditions that should be read into the model are defined by a list of keywords:

```
le_bound.types          : clim-isak clim-const clim-logan data
```

Each of the keywords defines a data set from which boundary conditions should be read. The tracers provided differ per set. If two sets contain data for the same tracer, the data corresponding to a keyword later in the list will replace the data read before.

The implementation and configuration differs per set. Each of the sets has its own module in the source code, for example:

```
le_bound_clim_emep.F90
le_bound_clim_isak.F90
le_bound_clim_logan.F90
le_bound_data.F90
:
```

The most general set is 'data'. The configuration of this set allows for input from time series of netCDF files. In future, all boundary conditions will be read through this configuration, but for the moment it is not used yet for 'older' climatological boundary conditions. The most important boundary conditions are those obtained from a global model (typically 3 hourly 3D fields), or a boundary run from LE (1 hourly resolution 3D fields) and these are all read using the 'data' configuration.

12.2 Configuration of 'data' boundary condition set

The processing of the 'data' boundary conditions follows a number of stages, and is illustrated in Figure 12.1.

- › At the bottom, the original boundary condition data (blue) is obtained from for example a global model. The time series could be irregular by resolution and format.
- › This data is read from the files and interpolated in time (green). Typically the code will store two original fields (for example valid for 00:00 and 03:00) and interpolate this linearly to a target time (00:30). At this stage, the data is also regridded to the model resolution, but the original levels are still kept.
- › In the next stage, also vertical regridding is applied (orange). This stage is separate because the vertical structure of the model is time dependent, and vertical regridding could therefore not be done directly after reading.
- › In the final stage the actual boundary concentration fields (red) are formed. Eventually those concentrations are formed from a linear combination of original fields, for example a weight sum of dust modes that together form the 'fine' fraction in the model.

The next sections illustrate step by step the configuration of these stages.

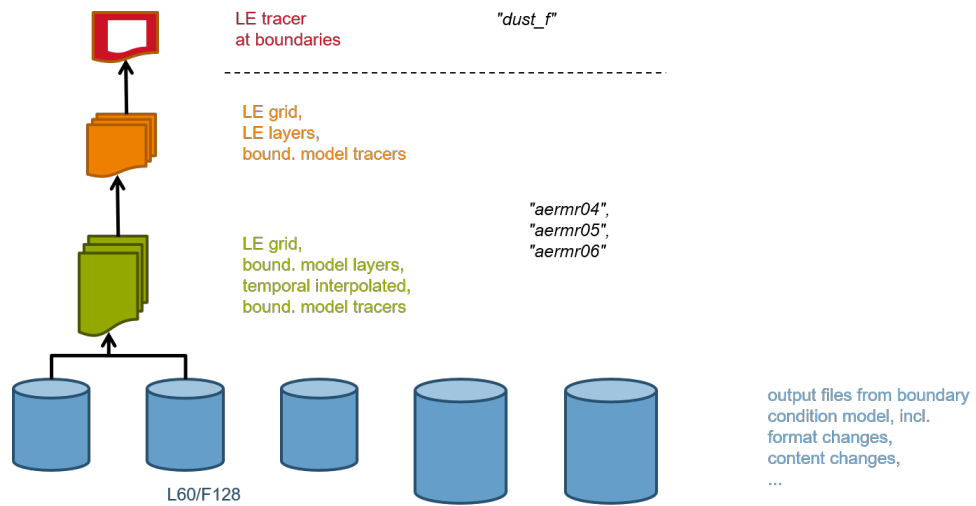


Figure 12.1: Data flow of boundary conditions.

12.2.1 Configuration for model variables

Figure 12.2 shows the top-level configuration of the boundary condition data. For each of the tracers in the model, the configuration should have a definition of how to form these concentrations from variables present in the boundary condition data. The example shows for example that the fine dust fraction should use a linear combination of the original dust modes.

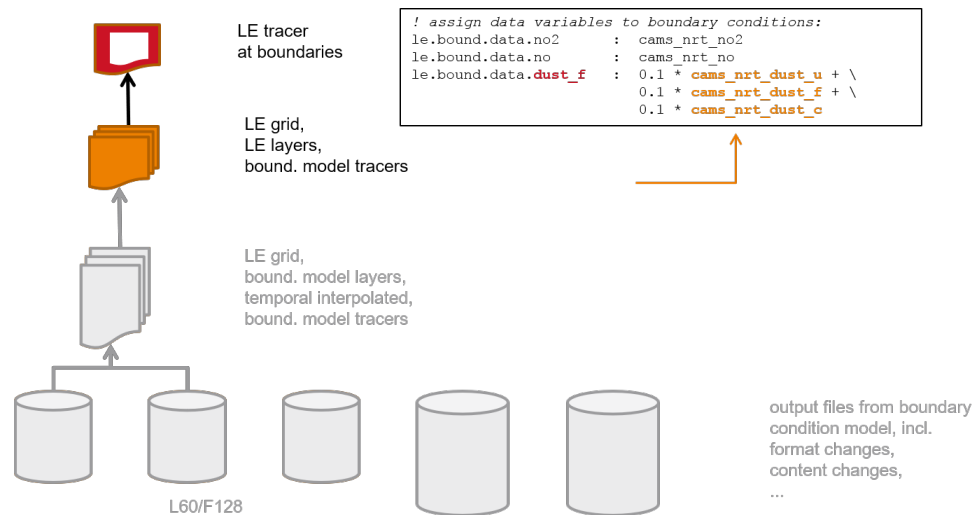


Figure 12.2: Top level configuration of boundary condition data.

12.2.2 Configuration of boundary condition variables

Figure 12.3 shows the configuration of the boundary condition variables. Each of these variables are stored in the 'LE_Data' structures that are also used for the meteorological data; see chapter 11 for a description.

In the example, the variable 'cams_nrt_dust_f' holds the fine dust mode read from the 'cams nrt' data set. As all boundary condition variables it holds concentrations on the model grid cells and model levels. The vertical regridding is defined using a 'call' description that tells the model to use an average over the original layers. This requires variables holding the original half level pressures and concentrations; also these should be present as 'LE_Data' variables. Here we simply added a double underscore to the names to define variables on the original levels:

```
__cams_nrt_hp
__cams_nrt_dust_f
```

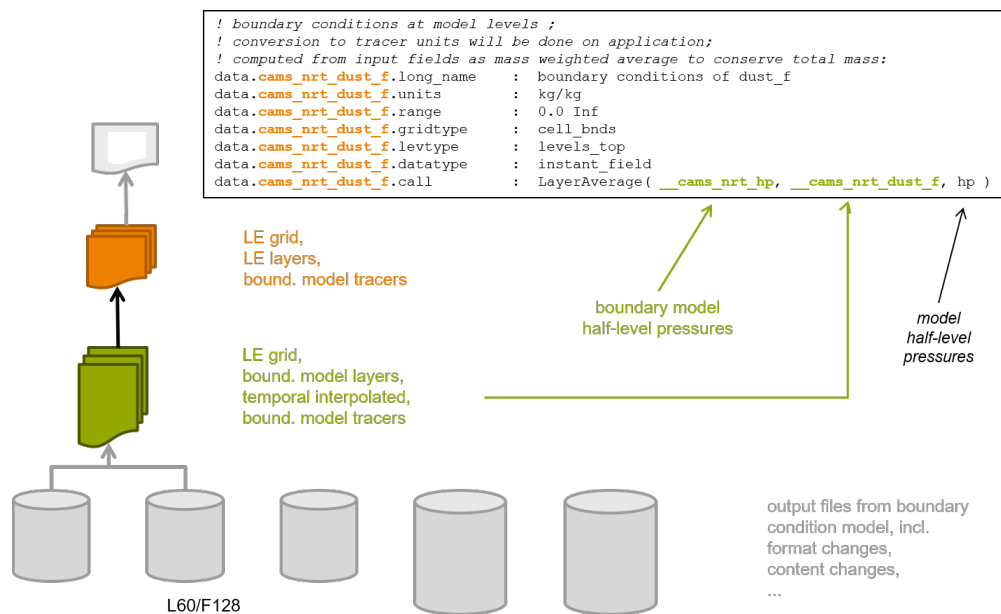


Figure 12.3: Configuration of regridded boundary condition variables.

12.2.3 Configuration of boundary condition variables on original levels

Figure 12.4 shows the configuration of the boundary condition variables on the original levels. Also these are configured as 'LE_Data' variables. In the vertical these variables are defined on input levels. The variables should be read from input files; an 'input' keyword is used for the configuration of a series of input files.

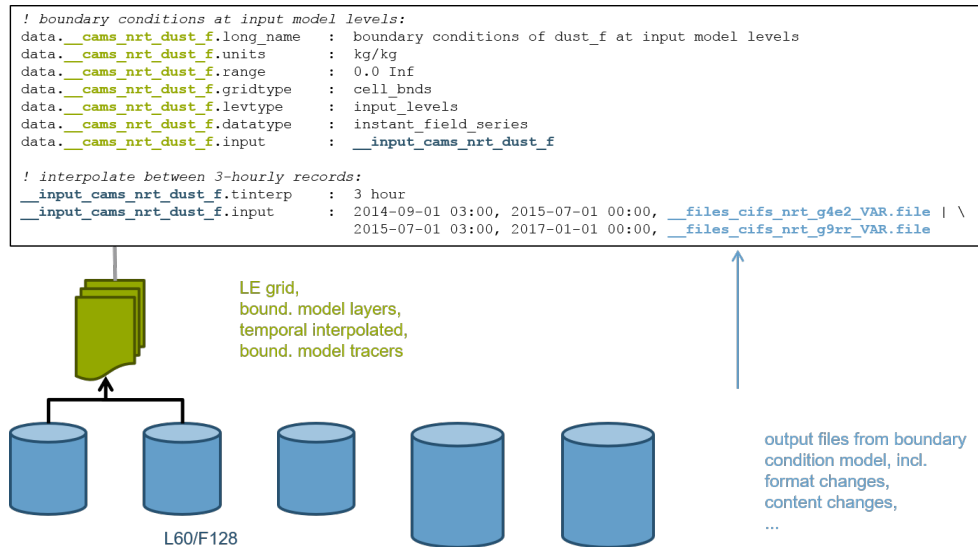


Figure 12.4: Configuration of boundary condition variables on original levels.

12.2.4 Configuration of boundary condition files

The final configuration is the description of the input file names and variables to be read. Figure 12.5 shows a part of this configuration for the dust variable.

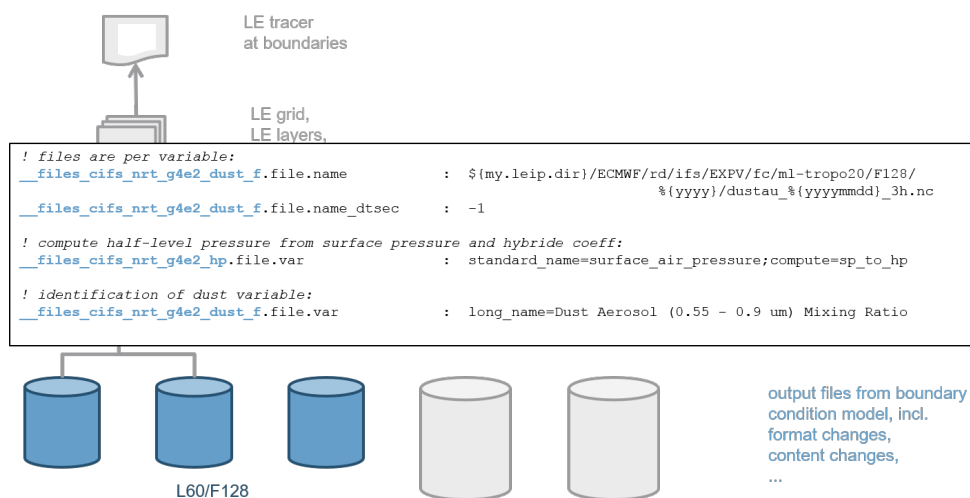


Figure 12.5: Configuration of boundary condition files and variables.

13 Land use maps and deposition parameters

13.1 Introduction

From LOTOS-EUROS v3.0.000, the land-use classification is based on a three-tiered land-use approach: climate zone, land use, and vegetation types as tier 1, 2 and 3, respectively. In this way, a coniferous tree in Norway can be distinguished from a coniferous tree in the Netherlands by their climate zones (continental and temperate, respectively), and it is possible to define model parameters specific to vegetation types. For instance, the deposition characteristics on potato plants may be very different from those on maize, despite the fact that they share the same land use class (arable land). See Manders and team [1] for details on this approach.

The three-tiered approach needs a land use map and a file with deposition parameters to perform calculations of dry-deposition fluxes of gases and particles. The land use map is a gridded map of fractions of all combinations of climate zone, land use class, and vegetation type. The deposition parameter file contains all land-use-dependent parameters for the dry deposition models for gases and particles. In this approach, the land use map is leading, i.e., each combination of the three tiers appearing in the map, should have a corresponding entry in the parameter file. For instance, a grid cell in the temperate climate zone on the map may contain arable land with maize on it. The deposition parameter file then should contain values for all parameters for the combination arable land/maize in the temperate climate zone. If not, the run will crash with an appropriate error message.

13.2 Data specification

The user needs to provide both the land use fraction map and the deposition parameters as NetCDF files. The format of these files is described in the next section. In the configuration file 'lotos-euros-expert.rc', the paths to these files have to be set:

```
le.landuse.input.file      : //path/to/landuse/fractionmap.nc
le.depac_data.file        : //path/to/deposition/parameterfile.nc
```

where obviously the pathnames need to correspond to your data files. A standard land-use fraction map and deposition parameter file are supplied with your OpenLE distribution. The land-use map was created by combining the Köppen-Geiger climate zone map with ESA2015 and Corine2018, as described in Manders and team [1].

13.3 File formats

13.3.1 Land-use fraction map

A NetCDF file should be supplied with a gridded map of fractions of vegetation types. Each grid cell covers a certain amount of land, which may belong to different climate zones and land use classes and contain different types of vegetation. The format is as follows.

```
netcdf tier123_example_map {
    dimensions:
        lu_types = 315 ;
        string11 = 11 ;
        longitude = 1380 ;
        latitude = 1560 ;
    variables:
        char lu_types(lu_types, string11) ;
        lu_types:long_name = "land-use types" ;
        lu_types:units = "1" ;
        float longitude(longitude) ;
        longitude:_FillValue = NaNf ;
        longitude:standard_name = "longitude" ;
        longitude:units = "degrees_east" ;
        float latitude(latitude) ;
        latitude:_FillValue = NaNf ;
        latitude:standard_name = "latitude" ;
        latitude:units = "degrees_north" ;
        float lu_fraction(lu_types, latitude, longitude) ;
        lu_fraction:_FillValue = NaNf ;
        lu_fraction:long_name = "land-use fraction" ;
        lu_fraction:units = "1" ;

        // global attributes:
        :Conventions = "CF-1.6" ;
}
```

The file has the following dimensions:

- › lu_types and string11 are the dimensions of the list of land use/vegetation combinations. string11 is always 11 characters long, since each LU/vegetation type is a combination of 3 three-letter abbreviations separated by underscores
- › longitude and latitude are the dimensions of the cell center coordinates

The variables are as follows.

- › lu_types(lu_types, string11): Abbreviation of the combination climate zone, land use class and vegetation type. For instance, the tag tmp_ara_whw represents the temperate climate zone, arable land as land use class and winter wheat as vegetation.
- › longitude(longitude). latitude(latitude): Cell center coordinates
- › lu_fraction(lu_types, latitude, longitude): The fraction of the cell at (latitude, longitude) that is taken up by the land use/vegetation combination lu_types.

Notes

- › the sum of all fractions over lu_types should equal 1 in each cell. If it is lower than 1, part of the surface area of the cell will not be available for deposition, if it is higher than 1, part of the surface area of the cell is counted double (i.e., there can be no overlap). Both situations lead to errors in the calculation of deposition.
- › the file should adhere to the NetCDF Climate and Forecast Metadata Conventions version 1.6 ([2], [3]).

13.3.2 Deposition parameter file

A NetCDF file should be supplied with all parameters for the dry deposition models for gases (DEPAC, Zanten et al. [4]) and particles [5]. Most of the parameters (as shown below) are a function of climate zone and land use/vegetation combination, a minor amount is also a function of the species that deposits. The format is as follows (abbreviated list for the sake of conciseness).

```
netcdf example_deposition_parameter_file {
    dimensions:
        climatezone = 7 ;
        landuse_vegetation = 69 ;
        species = 21 ;
        string3 = 3 ;
        string15 = 15 ;
        landuse_class = 13 ;
        string30 = 30 ;
        string7 = 7 ;
        string50 = 50 ;
        string10 = 10 ;
    variables:
        double A_bio_no(climatezone, landuse_vegetation) ;
        A_bio_no:_FillValue = NaN ;
        A_bio_no:long_name = "A_bio_no parameter for temperature dependence...
        ... of NO emissions per land use class" ;
        A_bio_no:units = "m" ;
        .
        .
        .
        int64 seasalt_flag(climatezone, landuse_vegetation) ;
        seasalt_flag:long_name = "Flag - to determine seasalt; 0:No seasalt...
        ...; 1:Can emit seasalt" ;
        seasalt_flag:units = "-" ;

        // global attributes:
        :author = "Leon Geers" ;
        :institution = "TNO" ;
        :title = "Land Use and Climate Dependent Deposition Parameters" ;
        :Conventions = "CF-1.6" ;
        :NCO = "netCDF Operators version 5.2.4 (Homepage = http://nco.sf....
        ...net, Code = http://github.com/nco/nco, Citation = 10.1016/j....
        ...envsoft.2008.03.004)" ;
        :history = "..." ;
}
```

The file has the following dimensions:

- › climatezone and string3 for the climate zone dependency of parameters. The latter dimension is always of size 3, since the climate zone dimension is a three lettered abbreviation. The climate zone names are max. 15 characters (dim. string15).
- › landuse_vegetation and string7 for the land use/vegetation dependency of parameters. Land use/vegetation combinations are represented with two three letter abbreviations separated by an underscore (e.g., ara_def for the default vegetation of arable land). The names of the land use/vegetation combinations are max. 50 characters (dim. string50).
- › species and string10 for species dependency of parameters. The species names each takes up max. 10 characters.

Table 13.1: Variables defined in the deposition parameter file.

Parameter	Dimensions	Description	Unit
landuse_class_names	landuse_class	The full name of the land use class	
climatezone_names	climatezone	The full name of the climate zone	
landuse_vegetation_names	landuse_vegetation	The full name of the land use/vegetation combination	
A_bio_no	climatezone, landuse_vegetation	A_bio_no parameter for temperature dependence of NO emissions per land use class	m
A_lu	climatezone, landuse_vegetation	A_lu parameter for Zhang model per land use class.	
alfa_lu	climatezone, landuse_vegetation	Alfa_lu parameter for Zhang model per land use class.	
alpha	climatezone, landuse_vegetation	Alpha value for light correction of stomatal resistance	m ² W ⁻¹
base_coeff_no	climatezone, landuse_vegetation	base_coeff_no parameter for temperature dependence of NO emissions	m
D_EGS	climatezone, landuse_vegetation	Shift of End of Growing Season	deg ⁻¹
D_SGS	climatezone, landuse_vegetation	Shift of Start of Growing Season	deg ⁻¹
Diffusivity	species	Diffusivity of species in air at 298K	m ² s ⁻¹
EGS_50	climatezone, landuse_vegetation	End of Growing Season at 50 deg lat	
ELEN	climatezone, landuse_vegetation	Length of End Phase of Growing Season	
fallow_flag	climatezone, landuse_vegetation	Flag - to determine fallow; 0:No fallow; 1:Fallow under circumstances; 2:Always fallow	
fmin	climatezone, landuse_vegetation	Minimum correction factor for stomatal resistance	
fst_th_Y	climatezone, landuse_vegetation	O3 stomatal flux threshold Y	nmol s ⁻¹ m ⁻²
gamma_lu	climatezone, landuse_vegetation	Gamma_lu parameter for Zhang model per land use class.	
gamma_soil_c_fac	climatezone, landuse_vegetation	Gamma correction factor for calculating soil compensation point	
gamma_soil_default	climatezone, landuse_vegetation	Gamma parameter for calculating soil compensation point	
gamma_stom	climatezone, landuse_vegetation	Gamma parameter for calculating stomata compensation point	
gs_max	climatezone, landuse_vegetation	Maximum leaf stomatal conductance for ozone	m s ⁻¹
ice_flag	climatezone, landuse_vegetation	Flag - to determine ice; 0:No ice; 1:Ice under circumstances; 2:Always ice	
ipar_snow	species	Parametrization type of Rc above snow	
LAI_max	climatezone, landuse_vegetation	Maximum Leaf Area Index	m ² m ⁻²
LAI_min	climatezone, landuse_vegetation	Minimum Leaf Area Index	m ² m ⁻²
Ld	climatezone, landuse_vegetation	Cross wind leaf dimension (Simpson, 2007) per land use class.	m
R_inc_b	climatezone, landuse_vegetation	In-canopy resistance parameter b	
R_soil	climatezone, landuse_vegetation, species	Soil resistance	s m ⁻¹
R_soilfrozen	species	Soil resistance for frozen soil	s m ⁻¹
R_soilwet	species	Soil resistance for wet soil	s m ⁻¹
SAI_a	climatezone, landuse_vegetation	SAI = SAI_a * LAI + SAI_b	
SAI_b	climatezone, landuse_vegetation	SAI = SAI_a * LAI + SAI_b	
SAI_b_dep	climatezone, landuse_vegetation	SAI dependent on LAI + Growing Season?	
SGS_50	climatezone, landuse_vegetation	Start of Growing Season at 50 deg lat	
SLen	climatezone, landuse_vegetation	Length of Starting Phase of Growing Season	
smi_level	climatezone, landuse_vegetation	Level for soil-moisture-index.	
temp_coeff_no	climatezone, landuse_vegetation	temp_coeff_no parameter for temperature dependence of NO emissions	m
Tmax	climatezone, landuse_vegetation	Maximum temperature for temperature correction of stomatal resistance	°C
Tmin	climatezone, landuse_vegetation	Minimum temperature for temperature correction of stomatal resistance	°C
Topt	climatezone, landuse_vegetation	Optimum temperature for temperature correction of stomatal resistance	°C
vpd_max	climatezone, landuse_vegetation	Maximum vapour pressure deficit	kPa
vpd_min	climatezone, landuse_vegetation	Minimum vapour pressure deficit	kPa
water_flag	climatezone, landuse_vegetation	Flag - to determine water; 0:No water; 1:Water under circumstances; 2:Always water	
z0_wind_dep	climatezone, landuse_vegetation	Roughness length for heat and trace gases dependent on wind?.	
z0dust_emis	climatezone, landuse_vegetation	Roughness length for dust emissions per land use class.	m
z0h_fallow	climatezone, landuse_vegetation	Roughness length for heat and trace gases on fallow land.	m
z0h_snow_surface	climatezone, landuse_vegetation	Roughness length for heat and trace gases on snow.	m
z0m_fallow	climatezone, landuse_vegetation	Roughness length for momentum on fallow land.	m
z0m_snow_surface	climatezone, landuse_vegetation	Roughness length for momentum on snow.	m
zcanopytop_fallow	climatezone, landuse_vegetation	Canopy top height per land use class on fallow land.	m
arable_flag	climatezone, landuse_vegetation	Flag - to determine arable; 0:No arable land; 1:Arable land	
seasalt_flag	climatezone, landuse_vegetation	Flag - to determine seasalt; 0:No seasalt; 1:Can emit seasalt	
desert_flag	climatezone, landuse_vegetation	Flag - to determine desert; 0:No desert area; 1:Desert area	
h	climatezone, landuse_vegetation	Height of surface elements	m

- landuse_class for the land use class names. The dimension is a three lettered abbreviation (dim. string3) and the land use class names themselves have a maximum number of 30 characters (string30).

The variables are given in Table 13.1. For each parameter, the dimensions are listed (the string length was not included for conciseness), a description is given and the units, where applicable. The description and units originate from the variable attributes of each parameter in the NetCDF file.

Notes

- For all climate zones and land use/vegetation combinations that are present in the land use

fraction map, there should be values for each parameter in the deposition parameter file as well. If any of them are missing, LOTOS-EUROS will crash. It is not a problem if there are land use/vegetation types represented in the deposition parameter table that do not have any fraction defined in the map. These will simply remain unused.

- › The format allows adding new land use classes and vegetation types.
 - Each land use class needs to have a default vegetation represented by the abbreviation 'def' in the dimension and 'Default' as the vegetation name. The values of each parameter for the default vegetation are the values representing an average for all vegetation in the land use class.
 - In case a new vegetation type is added, it should be classified under a specific land use class. Initially, values of parameters that are not available can be copied from the default vegetation type of that particular land use class.
- › Parameter values for specific climate zones and land use/vegetation combinations can be retrieved from literature. The values in the current file are mostly based on the defaults from DEPAC ([4]) and valid for Northwestern Europe. A table of some of the parameter values is given in the LOTOS-EUROS Reference Guide ([1]).
- › The file should adhere to the NetCDF Climate and Forecast Metadata Conventions version 1.6 ([2], [3]).

14 Generation of chemistry code

14.1 Overview

With the number of tracers growing throughout the years it became more and more difficult to manage the source files solving the chemistry. In addition, there was no easy way to quickly enable or disable some tracers or chemical reactions without having to change the source code.

It was therefore decided to let some of the tracer and chemistry related files be generated by the scripts, based on settings in the rcfile. The scripts used to generate the source files start with the name 'genes' (GENerate Sources), and this name is also used in the settings keywords.

The generated files are:

- › 'le_indices.inc', an include file with index parameters, names, units, and other settings for all tracers;
- › 'le_chem_work.F90', a module with routines that fill the reaction rates and perform a single step of the iteration step in used by the chemistry solver.

Default versions of the generated source files are included in the base source. Editing won't have any effect; instead change the rcfile settings or the tables that were used to generate them, as explained below.

The new source files are generated in the build directory. For debugging it might be useful to take a look at the generated files after creation; this is facilitated by the rcfile setting:

```
genes.show.command : nedit ${genes.files} &
```

In this case, the new files are loaded into the 'nedit' editor; the ampersand ensures that the editor keeps running as background process while the rest of the model setup continues.

14.2 Chemistry Scheme

LOTOS-EUROS includes chemistry schemes CBM4 and CBM7. Users can choose the chemistry scheme in the configuration file 'lotos-euros-expert.rc'.

```
! set genes.chemistry_scheme to cbm4, cbm7, or empty
genes.chemistry_scheme      : cbm4
```

The choice made here will determine which tracer and reaction tables are read in via switches in the 'lotos-euros-expert.rc'-file. In addition to the tracer and reaction tables, after CBM7 is chosen, the model reads in files with photolysis parameters and reaction types.

14.3 Tracer and chemistry properties

An important part of the configuration is done through so-called 'properties'. Table 14.1 shows some of the currently supported properties; their use will be explained later on. A list with supported properties is maintained in the 'lotos-euros-expert.rc'-file which is used to check the settings made by the user; see this expert rcfile for a complete list of all properties.

Tracer	chemical properties
cbm4	tracers/reactions of CBM4 scheme
cbm7	tracers/reactions of CBM7 scheme
sulphur	sulphur-only scheme (SO2 and SO4a, OH read in)
methane	methane-only scheme (CH4, OH read in)
co2	CO2 tracer
sf6	SF6 tracer
nmvoc	non-methane volatile organic carbon
radical	radical
ppm	primary particulate matter
ec	elementary carbon
pom	primary organic matter
sia	secondary inorganic aerosols
seasalt	sodium aerosols representing seasalt
dust	dust aerosols
m7	M7 aerosol scheme
vbs vbs_cg vbs_soa	for secondary organic aerosol modelling
basecation	base-cat-ion aerosols
hm	heavy metals
accum	accumulated species
biascorr	bias corrected species
aerosol	aerosol tracer
fine_mode	fine mode aerosol
coarse_mode	coarse mode aerosol
all_modes	total aerosol collecting all size modes
in-cloud	used to select implicit in-cloud chemistry reactions

Table 14.1: Tracer and chemistry properties.

To select the tracers and reactions that should be included in a simulation, a list with properties should be specified in the rcfile. Default selection in the main rc file is:

```
genes.chemistry_scheme      : cbm4
genes.group.selected        : ${genes.chemistry_scheme} sia ppm ec pom ...
...seasalt dust accum
```

This is used to select the appropriate lines from the tables described below.

14.4 Tracer table

The core of the tracer selection is a table with all supported tracers. A separate tracer table in .csv format is included in the model for CBM4 and CBM7 chemistry schemes.

```
base/000/data/tracers_cbm4.csv
base/000/data/tracers_cbm7.csv
```

The content looks like:

```
name , description , units , formula , properties , H-law , ... , rscale
NO , Nitric oxide , ppb , N + O , cbm4, 0.01, ... , 1
```

```

O3      , Ozone                        , ppb      , O 3      , cbm4, -2415.0, ..., 1
ALD     , Aldehyde                    , ppb      , C 2 + R, cbm4 nmvoc, 6300.0, ..., 1
PPM_f, prim.part.mat. fine mode, ug/m3, R      , ppm aerosol fine_mode, 0.0, ..., ...
...-999
:

```

For each supported tracer, the following values should be set:

- › *name* Short name, used in index variables i_O3 etc.
- › *description* Longer description, only used in comments in generated source files.
- › *units* Units of the tracer in the model; usually 'ppb' for gasses, and 'ug/m3' for aerosols.
- › *formula* Chemical formula (if possible). For some applications (labeling) it is useful to have at least the number of C, N, and S atoms in a molecule; for the remainder, use the symbol 'R'.
- › *properties* List with all tracer properties (table 14.1) that are applied to this tracer.
- › *H-law,T-fact,Reactivity,drydeftype,diffc,...,rscale* Tracer dependent parameters used for deposition calculation

To be enabled in the simulation, a tracer should have at least one of the selected properties.

14.5 Reaction table

14.5.1 CBM4

Chemical reactions for the CBM4 chemistry scheme are also specified in a text file. The default table is available as:

base/000/data/reactions_cbm4.csv

The content looks like:

```

label , reactants      , products      , rate expression      , properties
R1     , NO2                , NO + O3       , 1.0 x <NO2_SAPRC99>  , cbm4
R3     , O3+NO              , NO2           , 2.64 @ 1450          , cbm4
:
RH1f   , SO4a_f + N2O5, SO4a_f + 2*HNO3, rk_het(ireac_N2O5_NH4HSO4a_f) , cbm4 sia
:

```

For each reaction, the following values are specified:

- › *label*: A short label for the reaction, used in parameter names.
- › *reactions*: Tracers reacting with each other; tracer names following the tracer table.
- › *products*: Tracer products.
- › *rate expression*: Description of the reaction rate, see also Appendix A in the Reference Guide. See the comment in the top of the reaction table for how the expression is expanded. Some reaction rates are set by a special routine in the code, for example for the heterogeneous reaction 'RH1f' in the example lines above.
- › *properties*: List with all tracer properties (table 14.1) that that should be present to have this reaction enabled.

Reactions are only enabled if ALL of its properties are selected in the rcfile. Thus, reaction 'RH1f' of the above example is only enabled if both 'cbm4' and 'sia' tracers are selected.

14.5.2 CBM7

Chemical reactions for the CBM7 chemistry scheme are also specified in a text file. The default table is available as:

base/000/data/reactions_cbm7.csv

Relevant parts of this file are columns A:W, with two skipped rows. This content of the subsection looks like:

```
Number, (label) , (rate), (empty), Type, Order, 1, ... 12, Photolysis, ...
...groups
1 , NO2 = NO + O , NO + O3, 0.0063, ... , -1 , 1, 0, ... , 0.0063 , ...
...cbm7
2, O2 + O + M = O3 + M , 6.11E-34, ... , 3 , 3, 6E-34, ... , , ...
...cbm7
:
```

For each reaction, the following values are specified:

- › *Number*: Reaction number
- › *label*: Full reaction equation, with products on the left side and reactants on the right side
- › *rate*: Pre-calculated reaction rate (ignored)
- › *type*: Reaction type of the equation. An overview of the reaction types and their equations can be obtained from the *rate documentation* in

base/000/data/reactions_cbm7.xlsx

Reaction type -1 denotes a photolysis reaction.

- › *1..12*: Reaction parameters used, refer to Table 3-3b in *rate documentation* as stated above.
- › *Photolysis*: Parameter used in photolysis equation
- › *groups*: List with all tracer properties (table 14.1) that that should be present to have this reaction enabled.

Columns not mentioned here are not used in the model and are only left for reference.

Reactions are only enabled if ALL of its properties are selected in the rcfile. Thus, reaction 'RH1f' of the above example is only enabled if both 'cbm7' and 'sia' tracers are selected.

14.6 Specification of table files

The name of the tables file should be specified in the rcfile. The default setting in the rcfile for CBM4 chemistry scheme is:

```
genes.tracers.file : ../data/tracers_cbm4.csv
genes.reactions.file : ../data/reactions_cbm4.csv
```

And for CBM7 chemistry scheme is:

```
genes.tracers.file : ../data/tracers_cbm7.csv
genes.reactions.file : ../data/reactions_cbm7.csv
```

In this case, the tables are found in the 'data' sub-directory next to the 'src' sub-directory in the build directory, thus:

<rundir >/build /data/

These table files are copied from the base directory, and eventually replaced by project specific versions. Therefore, to test a new table, simply put it in the 'data' sub-directory of a project directory. Alternatively, specify an absolute path in the 'lotos-euros-expert.rc'-file settings.

14.7 Tracer indices and arrays

The information on the selected tracers is used to create the file 'le_indices.inc'. This file is included into 'le_indices.F90', which provides the module 'Indices'. Through this module, the user can access the generated entities; see the comment in top of 'le_incdices.F90' for their definition.

15 Labeling

15.1 rc-file

To use the labeling method switch, set the labeling flag to True

```
!
my.with.labeling      :  True
```

Define number of labels (default labels are added automatically)

```
!
labels.nlabel      :  2
```

Define which tracer to be labelled. Note that in a group, all tracers should be labelled

```
!
labels.labelled.specs :  ${N_tracers}  ${S_tracers}  ${unreactive_tracers}
```

15.2 Code

In module 'SA_Labeling.F90', the labels are defined hardcoded. Subroutine 'SA_Label_Definition' is use for this.

Step 1:

Define the names of the labels and define short names with maximum of 10 characters to get proper output for Grads-ctl files.

Note number of defined labels should be equal to number defined in rc-file. In the example, labels are defined for Countries (Netherlands and abroad). Also example lines for emission-sector labeling are provided.

```
!
SA_Label_Names(1) = 'NLD'
SA_Label_Names(2) = 'Abroad'
!SA_Label_Names(..) = '..'
```

Step 2:

Match incoming emissions to the label numbers. Each incoming emission must be attached to a label

Example for countries (with label on Netherlands and Abroad)

```
if ( icountry == -999 ) then
! undefined countries
def_label = 2
else if ( Emis_countries(iset)%emis_country_names(icountry) == 'NLD' ) then
def_label = 1
else
```

```
def_label = 2  
end if
```

Example for sectors (with label on Industry and others)

```
if ( Emis_Sectors(iset)%emis_sector_names(icat) == 'Industry' ) then  
def_label = 1  
else  
def_label = 2  
end if
```

15.3 Output

From the labeling module, extra outputfiles are generated. (labelled-conc-sfc, labelled-cond-3d). Properties of those files are identical to matching concentration files described in [Section 16](#), with an extra dimension 'label' to display the concentration resulting from each predefined label.

16 Model output

16.1 Output frequency

The user should specify the 'output' time-step in the rcfile:

```
! output time step in minutes :  
timestep.output      :    60
```

A typical value is 1 hour. The model will arrive at every multiple of this output-time-step and put out simulated values.

16.2 Gridded output

The output of gridded fields is controlled by the rcfile. The supported output types are described below. It is possible to put out files of the same type but with different properties, for example files with concentration fields at the surface for many tracers, and files with 3D fields for some selected tracers only.

The gridded output is written to NetCDF files. By default the files are structured following the CF-conventions. GrADS description files are added for visualization (see section [19.1](#)).

16.2.1 Concentration fields

For output of concentration fields the following properties could be set:

- › which tracers to be put out:
 - model tracers;
 - accumulated tracers: total PM10, total carbon, etc; see indices.F90 for the supported accumulation;
 - bias corrected tracers;
 - all-advected; all tracers which are advected in the model, used for nesting approach
- › vertical axis:
 - model levels, including the concentration at 2.5m (which is the measurement height, denoted as surface concentrations) and the top boundary layer;
 - heights relative to orography;
 - elevations relative to sea-level;
- › whether to put out the cell height too;
- › temporal resolution (typically fields are put out every hour);
- › collection per file: either daily or instantaneous;
- › horizontal coverage: by default the whole grid, but optionally:
 - bounding box to limit the horizontal area, for example to save storage space while producing boundary conditions for a zoom run;

- halo cells (boundary conditions values);

Template settings for output files with surface concentrations ('conc-sfc'), 3D fields at model levels ('conc-3d'), and 3D fields used as boundary conditions ('conc-bound') are available in the output rcfile.

16.2.2 Tracer total columns

Total columns could be put out for comparison with satellite observations. Note that this only includes the model layers now; the top boundary is not included yet, since usually there is no idea about its height. The following properties could be set:

- › which tracers to be put out (normal or accumulated);
- › temporal resolution (typically fields are put out every hour);
- › collection per file: either daily or instantaneous.

16.2.3 AOD columns

The Atmospheric-Optical-Depth is computed from the tracer concentrations and put out as columns.

16.2.4 Model data fields

For output of various other model data fields (e.g. meteorological fields), the following properties could be set:

- › which fields to be put out: meteo variables, stability fields, ...
- › vertical axis:
 - model levels, including the surface level and the top boundary layer which are for most fields a copy of the nearby model layer;
 - heights relative to orography;
 - elevations relative to sea-level;
- › temporal resolution (typically fields are put out every hour);
- › collection per file: either daily or instantaneous.

Template settings for output files with surface data ('meteo-2d') and with 3D fields at model levels ('meteo-3d') are available in the output rcfile.

16.2.5 Emission fields

The total emission for a certain tracer (or accumulated tracer) could be put out at regular times. The array that is put is 'emis_a' which contains for each tracer the total emission during the current (next) time step, independent of the source (anthropogenic, biogenic, sea-spray, etc). The following properties could be set:

- › for which tracers the emissions should be put out, including accumulated tracers;
- › vertical axis: either model layers or the total per grid cell;
- › temporal resolution (typically fields are put out every hour);
- › collection per file: either daily or instantaneous.

16.2.6 Dry and wet deposition

The dry- and/or wet-deposition budgets per output time step could be put out with the the following properties:

- › for which tracers the deposition should be put out, including accumulated tracers;
- › vertical axis: either model layers or the total per grid cell;
- › temporal resolution (typically fields are put out every hour);
- › collection per file: either daily or instantaneous.

16.2.7 Deposition parameters

Deposition variable such as resistances and deposition velocities could be put out with the following properties:

- › for which tracers the variables should be put out (if tracer dependent);
- › the landuse classes for which variables are valid;
- › vertical height, for example needed for dry deposition velocities.

16.2.8 Daily budgets

The daily budgets are updated every time step and are put out at midnight. The following budgets could be put out:

- › dry deposition flux of SO_x, NO_x, or NH_x;
- › wet deposition flux of SO_x, NO_x, or NH_x;
- › ozone dry deposition flux per landuse;
- › ozone daily maximum;
- › average NH₃ concentration in soil.

16.3 Satellite validation output

For special purposes also the following fields could be produced:

- › Simulation of OMI NO₂ columns.
- › Simulation of MODIS AOD columns.

The horizontal model grid is used to sample the satellite pixels. This output therefore requires that information on the satellite pixels is available.

16.4 Observation simulation

The standard method to produce simulations of concentrations at observation sites is to extract them from the (2D or 3D) gridded output.

An alternative is to use the MAORI (Model And Output Routine Interface) routines. The MAORI interface is configured via the rcfile by specification of a table file with station location and settings to define the simulated tracers etc. This is mainly used in the Kalman Filter context to simulate state values at observation sites; in the default model the code is present, but the use is not fully supported yet.

16.5 Emission summary

A summary of the emissions is written automatically to the output directory. Currently this is only supported for anthropogenic emissions read from files in TNO format. Since these are year-dependent, the summary is written for every year that the simulation covers. A summary consists of four comma-separated-value file (plain text): a list of the emission categories, a list with country codes and names, a table with total emissions for a component per country, and similar per country and emission category. The file is useful for verification of the emission input.

17 Restarting a run

When a run has crashed at some point due to some problem, the restart file can be used to restart the simulation at the day where it ended, without the need for spin-up. To use this option, change the date of the simulation in the main rc-file to the date of the last restart file. Thus, when the last restart file is:

```
LE_mysimulation_state_20120823_0000.nc
```

then the day and time for the simulation should be:

```
2012-08-23 00:00:00
```

Enable startup from a restart file by the setting:

```
le.restart      : T
```

and make sure that the path and runid of the restart file are defined correctly.

It is safer but not mandatory to move the existing output directory to a different path (e.g. `output2`). Be aware that the `.ct1` files for GRADS (see section [19.1](#)) are re-initiated so that they start now with the time stamp of the restart. This is easily modified in a text editor when one recombines the data from the original simulation and the restarted simulation.

18 Parallelization

LOTOS-EUROS has two options to employ multiple processors: domain decomposition using MPI, and OpenMP. These two do not exclude each other, but in practice the domain decomposition is preferred because it scales better when increasing the number of processors.

When running with MPI (domain decomposition) it is also possible to assign dedicated processor to copy input files from an archive (slow) to a local drive (fast); on some systems, this could strongly decrease the run time.

18.1 Domain decomposition

With domain decomposition, the model grid is divided into a number of sub-domains, and each sub-domain is assigned to another processor (figure 18.1). The processors use the MPI library to communicate between the domains, in particular to fill the boundary arrays using simulated concentrations from other processors.

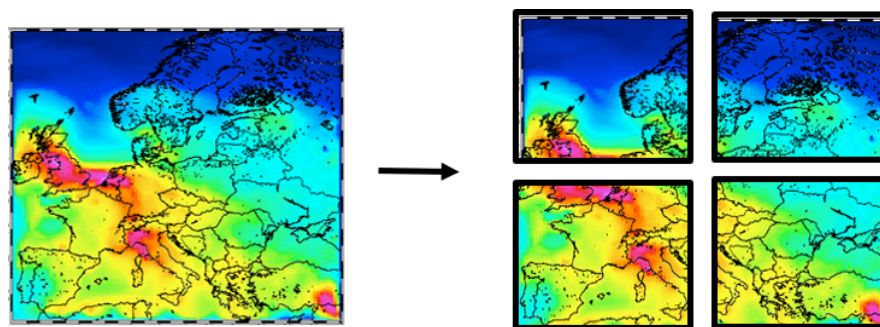


Figure 18.1: Illustration of domain decomposition.

18.1.1 Configuration

To run with domain decomposition, enable the MPI flag, specify the number of MPI tasks (sub domains), and specify how to decompose into sub-domains in x and y direction:

```
! enable MPI (True|False) ?
par.mpi                :   True

! number of mpi tasks:
par.ntask              :    4

! decomposition:
domains.x              :    2
domains.y              :    2
```

18.1.2 MPI compiler

The MPI library that is used for communication is linked with the model by using a special compiler wrapper around the native compiler. Popular MPI libraries are OpenMPI and MPICH, but also the Intel compiler suite usually comes with an MPI installation.

The name of the compiler wrapper that is used by the MPI library should be specified in the machine settings:

```
! compiler wrappers for MPI:
mpi.compiler.fc           : mpifort
```

18.1.3 Logging and error messages

Each sub-domain writes standard output to a separate log-file in the output directory. The domains are numbered sequentially starting from zero, and this number is part of the per-sub-domain log file name:

```
output/le-log-d00.out
le-log-d01.out
:
```

When an error occurs, it always happens first in just one of the subdomains. The error message is therefore in just one of the log-files only, since the processes handling the other sub domains are simply terminated. To see which processor received the error, check the content of the error file in the run directory:

```
---[lotos-euros_run.err]-----
MPI_ABORT was invoked on rank 3 in communicator MPI_COMM_WORLD
with errorcode 1.
...
```

In this example, the error was detected on processor 3.

18.2 I/O task

Reading input files (meteo, boundaries) could take a lot of time in case the file system is rather slow. On computing servers there is often a small but very fast disk available directly on the node, for example an SSD storage. When running with MPI (domain decomposition) it is possible to assign an extra processor to copy input files from an archive (slow) to the local drive (fast), which could strongly decrease the run time.

Figure 18.2 illustrates the allocation of processors to the model domains and to the i/o task. The root domain asks the i/o processor to:

- › check if selected input data for today is already available on the fast disk (here `/tmp`), and if not, copy it from the archive;
- › copy the data from the archive for tomorrow;
- › remove the data from yesterday from the fast disk, otherwise the fast disk might run out of space.

The copy and clean-up actions are performed by a script that is called from the i/o task. The script will install the input data for tomorrow on the fast disk, while the model is still simulating today; when the model arrives at tomorrow, the input data is already fast accessible.

To enable the i/o task, enable the following flag in the top-level settings:

```
! call installation script every day?
le.io.apply           : T
```

Using this flag, the machine specific settings should ensure that a job allocates an extra processor, such that the total number of MPI tasks is one more than the number of requested model sub-domains. For the TNO/HPC3 settings this is already configured.

The expert settings have configurations to ensure that the required input data for a standard run (meteo, boundary conditions, fire emissions) are installed on the fast disk. Here we shortly describe the most important settings that might require fine tuning in case of a non-standard simulations. The data sets to be copied temporary to the fast storage are identified by a list of keywords:

```
! list of data sets to be installed:
le.io.install.sets      : meteo gfas mc-bounds
```

In case of zoom run, also the boundary output from the outer simulation should be copied:

```
! list of data sets to be installed for zoom run:
le.io.install.sets      : meteo gfas mc-bounds le-bounds
```

For each of these datasets, define where to copy the data from (the archive) and where to copy it to; the values are taken from keys that should be defined in the machine specific settings:

```
! installation directory; empty for current:
le.io.install.meteo.dir : ${my.leip.dir}
! archive path with files to be installed:
le.io.install.meteo.arch : ${my.leip.arch}
```

Then provide a list of files that should be copied; for files including a time stamp the names should include templates for year/month/day/etc. For the meteo files this looks like:

```
! files to be installed:
le.io.install.meteo.files : ECMWF/od/ifs/0001/an/sfc/F1280/0000/*.nc \
ECMWF/od/ifs/0001/fc/sfc/F1280/%Y/*_%Y%m%d_1h.nc \
ECMWF/od/ifs/0001/fc/L137_CL42/F1280/%Y/*_%Y%m%d_3h.nc
```

18.3 OpenMP

Before the introduction of the domain decomposition, the LOTOS-EUROS model used OpenMP to employ multiple processors. The domain decomposition usually scales much better with

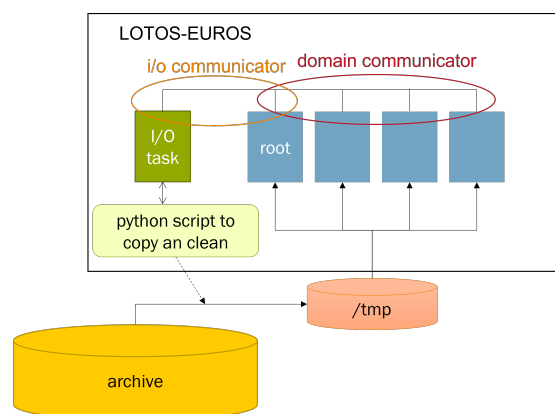


Figure 18.2: Allocation of processors for model domains and dedicated I/O task.

increased number of processors, but on some architectures it might still be beneficial to combine it with OpenMP.

18.3.1 Pragma's

The OpenMP standard uses special comments called *pragma's* in the source file to tell the compiler that multiple processors could be used for a certain piece of code:

```
!$OMP parallel
!$OMP do
do j = 1, ny
do i = 1, nx
! apply per grid cell:
call Something( c(i,j,:,:), ... )
end do ! i
end do ! j
!$OMP end do
!$OMP end parallel
```

18.3.2 Enable OpenMP

To run on multiple threads, enable the flag and specify the number of available threads:

```
! enable OpenMP (True, False) ?
par.openmp : True
! number of threads:
par.nthread : 0
```

It is our experience that usually 3 to 4 is a good choice, more than 8 will not lead to a further decrease of total runtime.

18.4 Timing

The time spent on different tasks in the model is continuously measured.

A timing profile is written to a text file with extension '.prf' in the output directory. In the header of this file, a pretty print of the absolute and relative time spent on a task and its sub-tasks is shown:

timer	system_clock	(%)
...		
model time loop	11504.66	
time step output	712.27 (6.2 %)	
time step setup	1223.58 (10.6 %)	
adjust	49.89 (0.4 %)	
advection	573.52 (5.0 %)	
chemistry	68110.19 (59.3 %)	
vertical diffusion	1144.94 (10.0 %)	
dry deposition	217.74 (1.9 %)	
sedimentation	24.93 (0.2 %)	
wet deposition	23.61 (0.2 %)	
emission	35.59 (0.3 %)	
other	679.41 (5.9 %)	
....		

19 Post processing and visualization

Post-processing and visualization of the model output is often very project and user specific. The netCDF file format compliant with conventions ensures that many packages can be used (e.g. Matlab, Python, IDL). In this chapter we give an overview of the available standard tools, which will help a user with the first steps.

19.1 GrADS

The *Grid Analysis and Display System* (**GrADS**) is an interactive desktop tool that is used for easy access, manipulation, and visualization of earth science data. It is very suitable to quickly browse through the data without scripting and perform a first screening of the output.

Note that GrADS cannot handle packed output.

19.1.1 Control files

LOTOS-EUROS supports the use of GrADS by adding control files (.ctl) to the output directories that are used by GrADS to read the output and define the correct grid, time range, etc.

19.1.2 Running

If GrADS is installed, use the following command to start:

```
grads
```

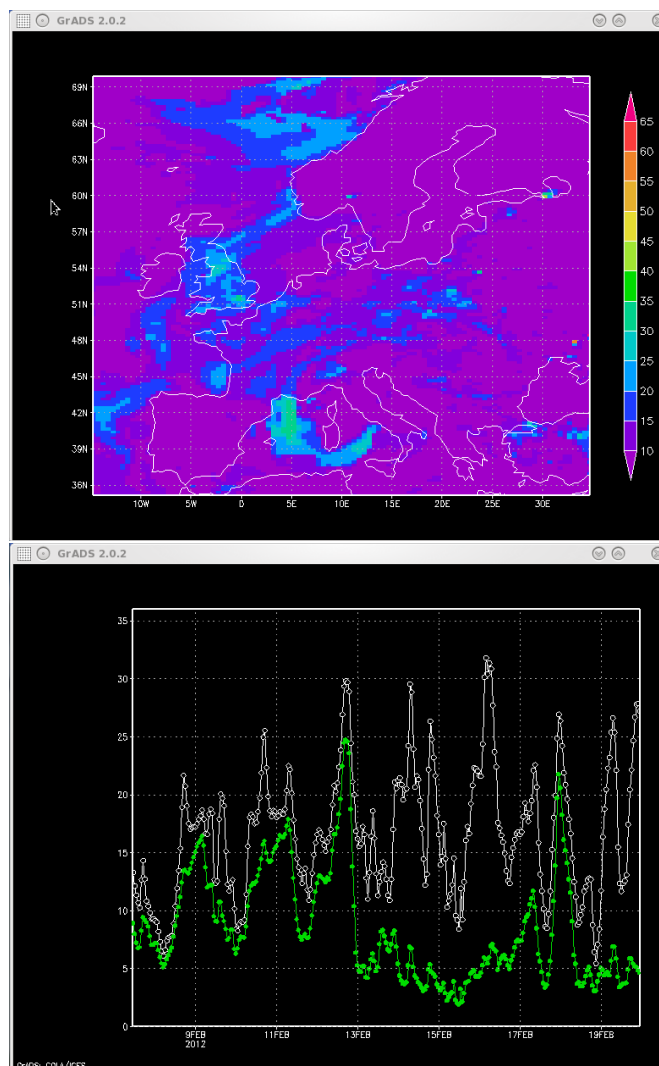


Figure 19.1: Screenshot of a GrADS graphics window with a snapshot of modelled PM10 concentrations and timeseries of PM10 and PM2.5 in gridcell corresponding to Utrecht (Netherlands)

20 Tools

The LOTOS-EUROS 'tools' are additional software packages next to the model.

For OpenLE, the tools are found in the 'tools' directory next to the base source (see section 4.2).

20.1 Create files with ECMWF meteorological data

Scripts to extract OpenLE meteorological data from the ECMWF archive, and to transfer the files to your local computer are present in:

`tools/meteo/`

An ECMWF member state account is required to run the scripts.

20.1.1 Extract data from ECMWF archive

The best (fastest) way to extract meteorological data from the ECMWF archive is to retrieve it from the MARS archive on the member state server ([ecgate](#)). After copying the 'tools/meteo' directory to `ecgate`, edit and run the following script to create data files:

```
./bin/OpenLE-meteo-ecmwf
```

This will first retrieve files from MARS in `grib` format, and then convert to `netcdf`. The standard `grib_to_netcdf` tool at `ecgate` does not provide completely CF-compliant files, and therefore small additional modifications are needed.

Variables that are originally accumulated from the start of a forecast are converted to averages over an interval. For example, precipitation fields are originally stored in 'mm water equivalent' since start, but converted to 'mm/s' average during a (3 hourly) time interval.

The data volume is reduced significantly by combining vertical layers. How to combine levels is defined in the header of the script; for OpenLE we created files with about 20 layers in the troposphere. To combine the levels a Fortran program is compiled that is present in the 'src' directory. Compilation settings are defined in:

```
rc/openle-meteo.rc
```

Extracting data could take a long time! Especially fields on ecmwf model levels (temperature, wind) are expensive to extract. The `ecgate` server does not allow jobs to run in foreground for longer than 30 minutes; longer jobs should be committed to the queue system. Use the following script to create a sequence of jobs that will be submitted one by one to the queue system:

```
./bin/OpenLE-meteo-ecmwf-jobs
```

Edit this file to set the time range and select the long/short jobs. To see which jobs are currently running on your account, use:

```
squeue -u $USER
```

The job-id number in the first column could be used to cancel a running job:

```
scancel <JOBID>
```

20.1.2 Transfer meteo files from ecgate to local computer

Meteo files produced at `ecgate` could be transferred to a local computer using the script:

```
./bin/OpenLE-meteo-get
```

The script uses the '[ECaccess](#)' tools to scan a source directory on `ecgate` and get the files to the local machine if not present yet. Edit the settings in the top of this script to change locations etc.

21 Coding conventions

How should new parts of code be written ? Some ideas.

- › Files contain either 1 module or 1 main program.
- › Model specific files have a prefix equal to the model name:

```
le_data.F90
le_process.F90
:
```

- › Modules have the same name as the source file:

```
module LE_Process
...
end module LE_Process
```

- › Module names (thus file names) should reflect the content:

```
! Data that are required for many different modules
module LE_Data

...
end module LE_Data
```

- › The tasks to be performed by a module could be distributed over a number of sub modules:

```
LE_Data                # top module

LE_Data_nc  LE_Data_hdf ...      # specific

LE_Data_Base          # shared entities
```

In this case, other model routines should access the entities only via the top module:

```
use LE_Data, only : LE_Data_Setup
use LE_Data, only : temper, humid, tsurf
```

- › Public routines in a module should start with the module name:

```
module LE_Data

private
public  ::  LE_Data_Setup
...

contains

subroutine LE_Data_Setup( t1, t2, status )
...
end subroutine LE_Data_Setup

...

end module LE_Data
```

- › If a module defines public data, a module initialization and finalization routine should be provided. These routines might be dummy to be prepared for future extensions.

```

module LE_Data

  private
  public  :: the_answer
  public  :: LE_Data_Init , LE_Data_Done
  ...

  integer      :: the_answer

  contains

  subroutine LE_Data_Init( status )
    ...
    ! something to be done:
    the_answer = 42
    ...
  end subroutine LE_Data_Init

  subroutine LE_Data_Done( status )
    ! nothing to be done.
  end subroutine LE_Data_Done

  ...

end module LE_Data

```

- › If a module defines a public type rather than public data, its name should start with the prefix 'T_' followed by the module name. An initialization and finalization routine should be created, eventually doing nothing:

```

module RcFile
  ...
  private
  public  :: T_RcFile , RcFILE_Init , RcFILE_Done
  ...
  type T_RcFile
    integer  :: id
    ...
  end type T_RcFile

  contains

  subroutine RcFILE_Init( rcf, fname, status )
    type(T_RcFile), intent(out)      :: rcf
    character(len=*), intent(in)     :: fname
    integer, intent(out)              :: status
    ...
    ! something to be done:
    rcf%id = 123
    open( unit=rcf%id, file=trim(fname), form='formatted', iostat=...
      ...status )
    ...
  end subroutine RcFILE_Init

  subroutine RcFILE_Done( rcf, status )
    type(T_RcFile), intent(inout)    :: rcf
    integer, intent(out)              :: status

```

```
...
! something to be done:
close( rcf%id, iostat=status )
...
end subroutine RcFile_Done

...

end type RcFile
```

- › Subroutines return an integer status value:

```
! return status:
!  <0  : warning
!   0   : ok
!  >0  : error

subroutine Process_Init( status )
integer, intent(out)    :: status
...
! ok
status = 0
end subroutine Process_Init
```

- › Functions are 'pure' and preferably 'elemental'.

References

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Signature

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