

# *Tutorial I. Installing the LMDZ model*



Ionela Musat / Laurent Fairhead

Laboratoire de Météorologie Dynamique

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[https://lmdz.lmd.jussieu.fr/pub/Training/Tutorials/Tutorial\\_1.pdf](https://lmdz.lmd.jussieu.fr/pub/Training/Tutorials/Tutorial_1.pdf)

# *Tutorial I. Installing the LMDZ model*



To install and run the LMDZ model you need to:

- 1/ **get source codes**: netcdf, IOIPSL, (XIOS\*), ORCHIDEE, **LMDZ**
- 2/ **compile** the codes
- 3/ **run a bench test** of 5 days

We have developped a script that does all three things:

**install\_lmdz.sh**

but there are 2 other ways

# *Tutorial I. Installing the LMDZ model*



There are **3 ways** to install LMDZ.

Right choice depends on the **machine** you are using, the **type of simulation** (**long** for research projects or **short** for development) you run and **the LMDZ version** you need.

## **1) using the `install_lmdz.sh` script** (⇒ this Tutorial)

- recommended method for **Linux PC or IDRIS**
- **development or short test run**
- **new LMDZ version available every 1-2 months**

## **2) using `modipsl` and `libIGCM`** ⇒ see IPSL Training course, 25-26 January or 1-2 February 2024)

- will install one of the configurations pre-defined by modipsl, for example **LMDZOR\_v6**
- recommended for **IDRIS, TGCC, CINES** and for **long simulations**, as it provides tested reference versions and scripts for launching and monitor long simulations.
- **new LMDZOR configuration available every 6-12 months**

## **3) by hand**

- recommended for people aiming **challenges (!)**
- you need to develop own scripts to get source codes for each component you need (**IOIPSL, ORCHIDEE, LMDZ**) and link them with the **netcdf library** installed on your machine.

# *Tutorial I. Using install\_lmdz.sh – Contents*



install\_lmdz.sh is using standard **shell tools and commands** ([wget](#), [gunzip](#), [tar](#), [gcc](#),...):

- Download the required codes archives
  - ↳ ancillary [libraries](#) ([netcdf](#), [modipsl](#), [IOIPSL](#), [XIOS\\*](#))
  - ↳ land surface model [ORCHIDEE\\*](#) (if requested)
  - ↳ [LMDZ](#) using [makelmdz\\_fcm](#) (or [makelmdz](#)) script
- Choose adequate [compiler options](#) and [build a Makefile](#)
- Run a [test bench](#)

Further details on [LMDZ version](#) (in French), in particular the main modifications between versions:

<http://lmdz.lmd.jussieu.fr/pub/LISMOI/trunk>

# Tutorial I. *install\_lmdz.sh* options



`./install_lmdz.sh -h`

<code>-v version</code>	LMDZ version YYMMDD.trunk	<code>20231022.trunk</code>
<code>-r release_nb</code>	LMDZ' svn release : <code>release_nb/last</code>	<code>----</code>
<code>-compiler compiler</code>	<code>gfortran / ifort/ pgf90/ mpif90</code>	<code>gfortran</code>
<code>-parallel mode</code>	parallelism type : <code>none/mpi_omp</code>	<code>none</code>
<code>-d grid_resolution</code>	grid resolution : <code>nlon x nlat x nlev</code>	<code>32x32x39</code>
<code>-bench</code>	launch or not a test bench : <code>1/0</code>	<code>1</code>
<code>-testing/unstable</code>	LMDZ' svn branch : <code>testing/unstable</code>	<code>testing</code>
<code>-name MODEL</code>	model folder <code>LMDZ\$version\$svn\$optim</code>	<code>LMDZ20231022.trunk</code>
<code>-netcdf PATH</code>	0, 1 or PATH to an existing netcdf	<code>1</code>
<code>-xios</code>	<code>with_xios="y"; need parallel=mpi_omp</code>	<code>----</code>
<code>-gprof</code>	compile with <code>-pg</code> to enable profiling	<code>----</code>
<code>-cosp</code>	run without, with COSP v1 or COSP v2: <code>NONE/v1 /v2</code>	<code>NONE</code>
<code>-rad</code>	RADIATIVE code : <code>oldrad/rrtm/ecrad</code>	<code>oldrad</code>
<code>-nofcm</code>	compile_with_fcm : <code>0/1</code>	<code>1</code>
<code>-SCM</code>	install 1D version automatically : <code>0/1</code>	<code>0</code>
<code>-debug</code>	compile everything in debug mode	<code>none</code>
<code>-opt_makelmdz</code>	call <code>makelmdz_fcm/makelmdz</code> with additional option	<code>none</code>
<code>-physiq</code>	physics' package to use : <code>physiq.def_\${physiq}</code>	<code>physiq.def_NPv6.1</code>
<code>-env_file</code>	<code>arch.env</code> environment file to overwrite the existing one	<code>----</code>
<code>-veget</code>	surface model to run : <code>NONE/CMIP6/xxxx</code>	<code>NONE</code>

# *Tutorial I. install\_lmdz.sh script*



**Download the script :**

```
 wget http://lmdz.lmd.jussieu.fr/pub/install_lmdz.sh
```

**Make it executable :**

```
 chmod +x install_lmdz.sh
```

**Execute the script, i.e. compile the model and run a 5-days test simulation**

```
 ./install_lmdz.sh -d 32x32x39 -name LMDZseq
```

=> modipsl.**20231022.trunk.tar.gz** => code sources

=> bench\_lmdz\_**32x32x39.tar.gz** => input files



## Principles:

Based on **makefiles** and **pre-processor** directives.

A **unique** procedure to compile the **different executables** (gcm, ce0l, 1d, ...)

An environment which allows the compilation of **different configurations** (zoom, resolution, physics, ...) from the same directory and source code.

An executable **compiled with ORCHIDEE** does not need to be **recompiled** to run without ORCHIDEE.

Pre-processor directives allow you to include specific parts of code in your main code. They are governed by what are called CPP keys (such as **CPP\_MPI**, **CPP\_COUPLE**, **CPP\_VEGET**, **INCA**) that are set by the installation script.

# Tutorial I. Code Compilation – the makefiles



Two different but similar scripts can be used to compile LMDZ :

- *makelmdz* : using the basic shell and our own scripts.
  - create the *dimensions.h* file using script *makdim* for the required resolution (this allows us to manage multiple resolution from the same directory)
  - create code dependencies with script *create\_make\_gcm*
  - create the *makefile*
  - define compilation and optimisation options
  - compile and creates the executable *gcm.e*
- *makelmdz\_fcm* : used by **MODIPSL**
  - creates the *dimensions.h file*
  - the *-arch* (needed) option determines the architecture of the target machine.  
Needed so as to read the right configuration file in the *LMDZ/arch* repertory
  - calls script *fcm* to generate dependencies and compile the code, creates an executable *gcm\_RESOLUTION\_PAR....e*

Simple example :  
`./makelmdz -d 48x32x11 -v false gcm`  
`./makelmdz_fcm -d 48x32x11 -v false gcm`

# *Tutorial I. Compilation : main options*



[makelmdz](#) main options :

[-h] :	help
[-d [[Imx]JMx]LM]:	IM, JM, LM are the x, y, z dimensions (def: 96x72x19)
[-p PHYS]:	to compile with libf/phyPHYS physics module, (def: lmd)
[-prod / -dev / -debug] :	to compile in production (default) / developpement / debug mode.
[-c false/MPI1/MPI2] :	ocean coupling : MPI1/MPI2/false (def: false)
[-v false/true] :	with or without vegetation (def: false)
[-chimie INCA/false] :	with or without INCA (def: false)
[-parallel none/mpi/omp/mpi_omp] :	parallelisation (default: none) : mpi, openmp or mix mpi_openmp
[-g GRI] :	grid definition in dyn3d/GRI_xy.h (def: regular)
[-io IO] :	choice of I/O library, left to the experts (def: ioipsl)
[-include INCLUDES] :	supplementary variables for includes
[-cpp CPP_KEY] :	supplementary CPP keys definition
[-filtre NOMFILTRE] :	use the filter in libf/NOMFILTRE (def: filtrez)
[-link LINKS] :	optional library links

[makelmdz\\_fcm](#) option:

-arch nom\_arch : name of target architecture

# *Tutorial I. Choosing which LMDZ version to work with*



Choose between the different available versions on the LMDZ web site:

<http://lmdz.lmd.jussieu.fr/pub/LISMOI.trunk>

Ask the LMDZ team for more information on which versions are actually used :

**lmdz-users@listes.ipsl.fr**

# *Tutorial I. What you need to run the LMDZ GCM (1)*



- Executable (LMDZ) file :
    - gcm.e
  - Parameters files
    - run.def, gcm.def, vert.def, physiq.def, traceur.def, config.def, etc
  - Start files
    - start.nc, startphy.nc
      - These files are created by the `ce0l.e` program or may be the result of previous runs
  - Boundary conditions file
    - limit.nc
      - Created by `ce0l.e`
  - Some optional input files **v** (depending on the simulation)
    - aerosols.nc, climoz\_LMDZ.nc, nudging input files (u.nc, v.nc,..), etc
- v** : these files have to be interpolated on the horizontal grid of the model

# Tutorial I. What you need to run the LMDZ GCM (2)



## I/O files for a LMDZ run

### Dyn/Phys/Boundary

start.nc  
startphy.nc  
limit.nc

### Parameters' files

run.def  
gcm.def  
vert.def  
physiq.def  
traceur.def  
orchidee.def\*  
guide.def

### LMDZ XIOS' files

iodef.xml  
context\_lmdz.xml  
field\_def\_lmdz.xml  
file\_def\_hist\*\_lmdz.xml

### ORCHIDEE XIOS' files

context\_orchidee.xml  
field\_def\_orchidee.xml  
file\_def\_orchidee.xml

LMDZ GCM  
gcm.e

ORCHIDEE

### Restart files

restart.nc  
restartphy.nc  
sechiba\_restart.nc

### Output files

histmth.nc	1mo
histday.nc	1d
histhf.nc	6h
histLES.nc	1h
histstn.nc	0.5h
histins.nc	1ts

### Optional files

climoz\_LMDZ.nc  
aerosols.nat.nc  
aerosols1980.nc  
tauswstrat.2D.nc  
taulwstrat.2D.nc  
solarforcing.nc  
sechiba\_start.nc

### ORCHIDEE input files

carteveg5km.nc  
cartepente2d.nc  
lai2D.nc  
PFTmap.nc  
routing.nc  
soils\_param.nc

# *Tutorial I. Running the model*



```
$> cd ~/LMDZseq/modipsl/modeles/LMDZ/BENCH32x32x39  
$> ls
```

gcm.e start.nc startphy.nc limit.nc config.def gcm.def orchidee.def physiq.def  
run.def traceur.def vert.def

```
$> ./gcm.e
```

or

```
$> ./gcm.e > listing 2>&1
```

To carry on a simulation that has been run, you have to copy the restart files obtained at the end of the previous run as new initial start files:

```
>$ mv restart.nc start.nc  
>$ mv restartphy.nc startphy.nc  
  
>$ ./gcm.e
```

# *Tutorial I. Has your run completed successfully ?*



**YES**

- ▶ you will then have a message saying ***Everything is cool*** on the standard output or in the output text file.
- ▶ The code will have created 2 restart files  
[restart.nc](#) and [restartphy.nc](#)  
needed to carry on your run
- ▶ and some output diagnostic files  
[histhf.nc](#), [histday.nc](#), [histmth.nc](#), etc  
to explore/view using ferret, grads, python...

# *Tutorial I. Has your run completed successfully ?*



**NO**

You must find out what the problem is.

Look for an error message in the output text file (called listing here), i.e. search for one of the following keywords/phrase:

***STOP, Houston, we have a problem, hgardfou, negative surface pressure, ...***

**Different typical errors :**

- **Technical problem** : a missing input file, an error in one of the \*.def or .xml file
- **Problem with the model's stability.**
  - instability in the physics are likely to be detected by **hgardfou**, which checks the model temperature has realistic values.
  - instability in the dynamics most often end up the run with a **negative surface pressure** (**integrd** routine) error message.
- ▶ **In any of these cases you will most probably have to adjust some flags in the .def files (modify dynamic' timestep and teta\* flags for a new resolution, for example).**
- **Otherwise you may have made some source code modifications that might not have been thoroughly tested or validated.**

# *Tutorial I. Installing the LMDZ model: Take-off infos*



- **Install LMDZ** using [install\\_Lmdz.sh](#)

```
wget http://lmdz.lmd.jussieu.fr/pub/install_Lmdz.sh  
chmod +x install_Lmdz.sh  
./install_Lmdz.sh -d 32x32x39 -name LMDZseq
```

- **Re-compile** and **re-run** an LMDZ simulation : [compile.sh](#) and [bench.sh](#)

```
cd ~/LMDZseq/modipsl/modeles/LMDZ  
./compile.sh  
cd BENCH32x32x39  
./bench.sh
```

- **LMDZ releases** : ask to [lmdz-users@listes.ipsl.fr](mailto:lmdz-users@listes.ipsl.fr) or look here

<http://lmdz.lmd.jussieu.fr/pub/LISMOI.trunk>