# Hands on tutorial session 1: First steps with the model

#### The LMDZ team

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This first tutorial focuses on installing and making basic first runs using the LMDZ model. This document can be downloaded as a pdf file:

wget http://www.lmd.jussieu.fr/~lmdz/Distrib/TD1\_en.pdf

which should ease any copy/paste of command lines to issue.

# **1** Prerequisits

To run LMDZ, you will need a significant amount of memory, so first ensure this is true. You can use the following command line (that you can also, for convenience, put in your /.bashrc file):

```
ulimit -Ss unlimited
```

If you are working on a laptop provided by the LMD and dedicated to these training sessions, you have to log in as the default (and sole) user util1 (the associated password will be given during the training session). On those machines, a working folder, named LMDZ, contains all the required archives ; if it's not already there, you can create it:

mkdir LMDZ cd LMDZ

You may want to copy this full folder on a USB key at the end of the sessions.

# 2 Running the install\_lmdz.sh script

The first step consists in downloading it from the LMD website and *blindly* running it (after having first set the access permissions to make it executable):

```
wget http://www.lmd.jussieu.fr/~lmdz/Distrib/install_lmdz.sh
chmod +x install_lmdz.sh
./install_lmdz.sh -d 48x36x39
```

Compilation takes around five minutes. Then, a test bench is downloaded (if not already on the disk) and a 1 day long test simulation on a regular  $48 \times 36$ -L39 grid is run ; messages about various downloads (via **wget**) and/or compiler informations are displayed.

The script should then run smoothly (if it isn't the case, immediately ask for some assistance) and end with messages of the likes of:

You can take advantage of the installation time to open a second terminal window and explore the downloaded directories and files.

install\_lmdz.sh will check if some archives (LMDZ, NetCDF library, etc.) are on the disk (in the ~/LMDZ/Distrib folder) and if not, will try to get them through the network using wget command. It will create the LMDZtesting directory ; inside, you will find subdirectories modipsl, which contains the model, and netcdf-4.0.1, which contains the NetCDF library.

In modipsl, you will find directory modeles, containing the LMDZ5 directory.

Once the test bench simulation has been launched (the final step of the **install\_lmdz.sh** script), you will also find a **LMDZ5/BENCH48x36x39/** directory from where you will be able to list the outputs of the run (even if the simulation is still running: it indeed takes a few minutes to complete the 1 day-long run on a single processor). Check out the contents of this directory and use your favorite software (Grads, Ferret,...) to browse the contents of the **histday.nc** file.

The default behavior of the install\_lmdz.sh script is to install the model using the gfortran compiler, which is fine on the laptops provided for this training course, but on some local machines, an old version of the compiler only is available, so the script stops with a message that invites you to switch to another compiler. In any case, you can change from compiler by editing the script.

Several other features of the script can be modified or unactivated just by editing few parameters in the file or specifying the appropriate option (run ./install\_lmdz.sh -h to learn about these). For example, you can disable the NetCDF download and installation (first operation performed by install\_lmdz.sh) in case this library is already present on the computer you are using. The same kind of adjustments is possible for ORCHIDEE and LMDZ <sup>1</sup>.

# 3 Changing the outputs

The model outputs are controled in file **config.def** by the following lines:

phys_out_filekeys=	у	У	n	У	n
phys_out_filenames=	$\verb+histmth+$	histday	histhf	histins	histLES
phys_out_filetimesteps=	5day	1day	1hr	6hr	6hr
phys_out_filelevels=	10	5	0	4	4
phys_out_filetypes=	ave(X)	ave(X)	ave(X)	<pre>inst(X) ;</pre>	inst(X)

You will obtain 3 files (apart from ORCHIDEE related outputs): histmth.nc, histday.nc and histhf.nc, containing respectively 0 days (should contain 1 record for the whole run, but the output frequency set for histmth.nc above is 5day, whereas the run was only 1 day long), daily and hourly averages.

You can find out which variables have been written in a given file by running the command:

#### ncdump -h histday.nc | grep long\_name

To get a histmth.nc with data, you'll need to run the model over 5 days. This means you have to change the value of nday in file run.def to 5.

You may also re-run the simulation with a higher output frequency for few variables (e.g. atmospheric temperature at 2m above the surface: **t2m**, surface pressure: **psol**, precipitation: **precip**, etc.), and other fields types (for instance instantaneous fields) by modifying **config.def**:

phys_out_filekeys=	у	у	n	У	n
phys_out_filenames=	$\verb+histmth+$	histday	histhf	histts	histLES
phys_out_filetimesteps=	5day	1day	1hr	1TS	6hr
phys_out_filelevels=	10	5	0	4	4

<sup>&</sup>lt;sup>1</sup>The LMDZ compilation embedded in **install\_lmdz.sh** uses the **makelmdz\_fcm** script rather than the default **makelmdz** script. To change this behaviour in the **install\_lmdz.sh** script find the instruction line **compile\_with\_fcm=0** and change it to **compile\_with\_fcm=1**.

phys_out_filetypes=	ave(X)	ave(X)	ave(X)	inst(X)	inst(X)
flag_t2m=	10	10	5	5	5
flag_psol=	10	10	5	5	4
flag_precip=	10	10	4	5	5

Note that unlike other frequencies, 1TS (1 Time Step) is written in capital letters. You can check in particular the **histts.nc** file and explain the special results.

### 4 Making some sensitivity test runs

You can now move on to changing parameters in a .def file. One may for instance deactivate the subgrid orography parametrizations by changing the **ok\_orodr** and **ok\_orolf** flags in **physiq.def**. One could also choose to change the value of the cloud water to rain conversion factor **cld\_tau\_lsc** (in **physiq.def**), or the atmospheric CO<sub>2</sub> concentration, etc. (ask around to find out the meaning of the various available parameters).

Just run the simulation (in a different directory to avoid overwritting output files) and investigate the differences between simulations (using diff or your favorite visualization software).

# 5 Switching to a different version of the model

The recommended LMDZ version is the latest from the *testing* branch, but the install\_lmdz.sh script you have launched is set to download the latest version of the *trunk* branch of the model, containing recent developments some of you may need. Usually, it is safer to use the *testing* branch, much more extensively tested than the *trunk* branch. Your version is dated November 30th 2016, which corresponds to *svn release* 2720. This information can moreover be obtained by checking at the last line of the file:

#### http://www.lmd.jussieu.fr/~lmdz/Distrib/LISMOI.testing

or directly from within the LMDZ directory:

```
cd ~/LMDZ/LMDZtesting/modipsl/modeles/LMDZ5 svn info
```

Note: you may encounter the following error when typing *svn info*:

```
svn: E155036: Voir la commande 'svn upgrade'
svn: E155036: The working copy at '/home/util1/LMDZ/LMDZtesting/modipsl/modeles/LMDZ5'
is too old (format 10) to work with client version '1.8.8 (r1568071)' (expects format 31).
```

If so, simply follow the suggested cure. Enter: svn upgrade. Then try again svn info. Similarly, for *trunk* versions, this information is available at the end of the file

http://www.lmd.jussieu.fr/~lmdz/Distrib/LISMOI.trunk

Alternatively, same information is available on the following web page, dedicated to this branch:

```
http://lmdz.lmd.jussieu.fr/utilisateurs/distribution-du-modele/
versions-intermediaires/versions-intermediaires-ou-testing-en?set_language=en
```

Using svn, it is possible to change from model version and select and older or newer svn release. To change from current version to *svn release* 2669 (a previous version), go to directory LMDZ5 and issue command:

svn update --revision 2669

Important: as you change svn revision, a conflict on a file (**bld.cfg**) will occur, answer tc ('tc' for theirs-full) to tell svn you want to use the default version and not your own modifications<sup>2</sup> of the file.

From LMDZ5 directory, run<sup>3</sup> ./makelmdz\_fcm -arch local -j 8 -rrtm true -d 48x36x39 -v orchidee2.0 -cpp ORCHIDEE\_NOZ0H gcm

(or alternatively ./makelmdz -arch local -d 48x36x39 -rrtm true -d 48x36x39 -v orchidee2.0 -cpp ORCHIDEE\_NOZ0H gcm).

Option -v orchidee2.0 -cpp ORCHIDEE\_NOZOH is not mandatory. It only indicates that we compile with Orchidee, but in fact, apart from the first bench, automatically launched by the install\_lmdz.sh script, it won't be used in this tutorial. Only routines modified by the svn update or depending on them will be recompiled.

Once the model has been successfully recompiled, run a new simulation. To do so, create a new subdirectory in LMDZ5 and copy boundary conditions, initial conditions and parameters files (limit.nc, start\*nc, \*.def) over from directory BENCH48x36x39, along with the newly created gcm executable.

If you used **makelmdz**, it will be **LMDZ5/gcm.e**.

If you used  $makelmdz_fcm$ , it will be  $LMDZ5/bin/gcm_48x36x39_phylmd_seq_orch.e$ . Then launch the run<sup>4</sup>:

./gcm.e > listing 2>&1

Redirection of the model outputs in a text file is usually the best way to keep a trace of the run.

Once the simulation is finished, you can compare, using the **diff** command and visualisation software, the simulation results (compare files **listing**, **hist\*.nc restart.nc** and **restartphy.nc** from the two simulations) which should be identical, except for some differences in the text outputs.

Once this test finished, revert to the latest *trunk* version of the model by going to directory LMDZ5 and using:

svn update

Note that using **svn update** without any specific revision number implies updating to the latest version on the branch (which is something you should do regularly if you want to keep up with model updates). Don't forget to recompile the model after any svn update !

### 6 Running an aquaplanet configuration

The simulations run in the previous example included initial and boundary conditions (**start\*.nc** and **limit.nc** files). It is also possible to run the model in "aquaplanet" configuration, with idealized initial and boundary conditions are used (no topography and imposed surface temperatures).

Make an **AQUAPLANET** directory where you will run the model. Copy over the **gcm.e** and **\*.def** files from another simulation to this directory. Edit the **gcm.def** file to set

read\_start=n
iflag\_phys=101

Note that there are various possibilities for flag **iflag\_phys**, values between 101 and 114 correspond to different choices of imposed SSTs (see routines **iniaqua** and **profil\_sst** in file **phyaqua\_mod.F90** in the **libf/phylmd** directory).

<sup>&</sup>lt;sup>2</sup>This conflict comes from the fact that the installation script *install\_lmdz.sh* adds a line in the **bld.cfg** script and that there have also been some (harmless) changes of these very same lines between svn revisions of that file. Thus svn cannot know which modifications should overrule and thus asks you.

 $<sup>^{3}</sup>$ To avoid typing such a long line over and over, it is more convenient to have it in a script, e.g. **compile.sh** to execute each time a compilation is required.

<sup>&</sup>lt;sup>4</sup>As for the compilation step, this step can be put in a script, e.g. **bench.sh**, as is the case in **BENCH48x36x39** 

Then run the model:

./gcm.e > listing 2>&1

Inspect the various outputs (e.g. zonal averages and deviations of meteorological quantities such as surface pressure *psol*) and their temporal evolution, and compare to previously obtained simulations. Experiment changing the length of the run (variable **nday** in the **run.def** file to investigate the evolution of the system from initial settings to a "converged" state.

If time permits, experiment running at a different resolution (e.g.  $96 \times 95$ -L39; which means you will have to recompile the model). Note that the number of time steps per day depends on the model resolution, and has to be higher for this resolution than it was strictly required for the  $48 \times 36$ -L39 case. The gcm.def file you've used has already **day\_step=480**, which is enough, but you can check that the model will crash if you reduce it significantly (try it !).

# 7 Preparing things for tomorrow's course

Tomorrow you will need to run the model (at  $48 \times 36$ -L39 resolution) and investigate the outputs. To prepare things for that you must, in a dedicated directory, run the model for 5 days, using the files provided in the **bench\_lmdz\_48x36x39.tar.gz** (only the run lenght, **nday=5** needs be changed).